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Expression of human serum albumin in pichia pastoris.

② A novel expression cassette, vectors and process for the secretion of HSA in Pichia pastoris cells. In accordance with the present invention there has been discovered an improved expression cassette for the production of HSA in Pichia pastoris comprising

a) a Pichia pastoris 5" regulatory region having a 5' end and a 3' end selected from the group consisting of the Pichia pastoris AOX1 5' regulatory region and the Pichia pastoris DAS1 5' regulatory region wherein the 3' end of the 5' regulatory region is operably linked to:

b) an HSA structural gene encoding a signal sequence and mature protein having a 5' end and a 3' end wherein the HSA structural gene has an ATG start codon within about 11 deoxyribonucleotides of the 5' end of said HSA structural gene; and operably inked to

c) a 3' termination sequence.

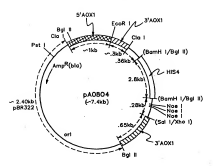


FIG. 1

Field of the Invention

This invention relates to the field of recombinant DNA biotechnology. In one aspect, this invention relates to a process for the expression of human serium albumin (HSA) in *Pichia pastoris*.

Background

Human serum alburnin is the most abundant plasma protein of adults. The concentration of alburnin is 40 mg/ml, or 180g of alburnin circulating throughout the human body for a 70 kg adult male. This protein maintains osmotic pressure and functions in the binding and transport of copper, nickel, calcium (weakly, at 2-3 binding) sites), bilirubin and protoporphyrin, long-chain fatty acids, prostaglandins, steroid hormones (weak binding) with these hormones promotes their transfer across the membranes), thyroxine, triidothyronine, and glutathione. According to Peters, T. and Reod, R.G. in Alburnin: Structure, Biosynthesia and Function, (Peters, T. and Sjoholm, J. eds.) 1977 p.11-20, over 10,000 kilograms of purified alburnin ats administered annually in the United States alone to patients with circulatory failure or with alburnin deceletion.

Currently the only commercial source of HSA is from fractionated blood. Considering the possible dangers of blood borne contaminants and pathogens, it would be a considerable contribution to the commercial production of HSA to develop alternate methods of producing HSA. With the advent of recombinant DNA technology, it is now possible to produce HSA by alternate methods.

HSA has also been expressed in Saccharomyces cerevisiae as disclosed by Etcheverry et al. in Biohechnology, August 1986, p. 726 and Arjum Singh in EPA 125,944. Eicheverry disclosed HSA expression intracellularly in a concentration of approximately 6 mg/l and the presence of cell-associated HSA. Hayasuke et al. also disclosed the expression of HSA in Saccharomyces' cerevisiae in combination as with the GAL 1 promoter and a signal sequence. Hayasuke et al. alsopast to have been able to achieve a secreted production level of 160 mg/l. As described in EPA344,459, HSA has also been expressed in Pichia pastoris as intracellular or cell-associated protein. Although the expression of HSA in yeast-cells, such as Saccharomyces cerevisiae and Pichia pastoris, is a significant step toward providing safe alternative sources of HSA, the expression of HSA as an intracellular or cell-associated protein is not desirable. Expensive and time consuming measures are required to recover and purify intracellular or cell-associated this Archiver of the production of the providing safe achieved to improve the yields and lower the production costs.

Thus, it would be a significant contribution to the art to provide yeast strains which secrete high levels of HSA.

It also would be a significant contribution to the art to develop a process which produces HSA in a manner that is easy to recover and purify.

Therefore, it is an object of this invention to provide strains which produce high levels of HSA.

It is a further object of this invention to provide a process which produces HSA in a manner that is easy to recover and purify.

Other objects and advantages of the present invention will be apparent to those skilled in the art from the present specification.

Summary of the Invention

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In accordance with the present invention there has been discovered an improved expression cassette for the production of HSA in Pichia pastoris comprising

 a) a 5' Pichia pastoris regulatory region having a 5' end and a 3' end selected from the group consisting of the Pichia pastoris AOX1 regulatory region and the Pichia pastoris DAS1 regulatory region wherein the 3' end of the regulatory region is operably linked to

b) an HSA structural gene encoding a signal sequence and mature protein having a 5' end and a 3' end wherein the HSA structural gene has an ATG start codon within about 6 deoxyribonucleotides of the 5' end of said HSA structural gene; and operably linked to

c) a 3' termination sequence.

in accordance with the present invention there has also been discovered Pichia pastoris strains transformed with an improved expression cassette for the production of HSA in Pichia pastoris comprising a) a 5 Pichia pastoris regulatory region having a 5' end and a 3' end selected from the group consisting of the Pichia pastoris AOX1 regulatory region and the Pichia pastoris DAS1 regulatory region wherein the 3' end of the regulatory region of poerably linked to provide the picking pastoris DAS1 regulatory region in the 1' end of the regulatory region in the 1' end of the regulatory region in the picking pastoris DAS1 regulatory region in the 1' end of the regulatory region is poerably linked to the provide pastoris that the picking pa

b) an HSA structural gene encoding a signal sequence and mature protein having a 5° end and a 3° end wherein the HSA structural gene has an ATG start codon within about 6 deoxyribonucleotides of the 5° end of said HSA structural gene; and operably linked to

c) a 3' termination sequence.

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In a further embodiment of the present invention, there has also been discovered a process for the secretion of HSA from transformed *Pichia pastoris* cells comprising

a) transforming Pichia pastoris with at least one vector having at least one expression cassette comprising

i) a 5' Pichia pastoris regulatory region having a 5' end and a 3' end selected from the group consisting of the Pichia pastoris AOX1 regulatory region and the Pichia pastoris DAS1 regulatory region wherein the 3' end of the regulatory region is operably linked to

iii) an HSA structural gene encoding a signal sequence and mature protein having a 5' end and a 3' end wherein the HSA structural gene has an ATG start codon within about 6 deoxyribonucleotides of the 5' end of said HSA structural gene; and operably linked to

iii) a 3' termination sequence; and then

b) culturing the resulting transformed *Pichia pastoris* under suitable conditions to obtain the secretion of HSA.

Detailed Description of the Figures

Figure 1 provides a representation of plasmid pA0804 which contains a linear site-specific integrative vector in the fragment clockwise from Bglll to Bglll. The structural gene may be inserted in the unique EccRl site of this plasmid. This plasmid may be recovered from the plasmid DNA of NRRL B-18114 by EccRl digest and gel electrophoresis to recover a linear -7.4 kb EccRl fragment corresponding to Figure 1.

Figure 2 provides a representation of pHSA13 in circular form.

Figure 3 provides a restriction map of the AOX1 5' regulatory region isolated from Pichia pastoris.

Figure 4 provides a restriction map of the DAS1 5' regulatory region isolated from Pichia pastoris.

Figure 5 provides a restriction map of the AOX1 3' termination sequence isolated from *Pichia pastoris*. Figure 6 provides a restriction map of the DAS1 3' termination sequence isolated from *Pichia pastoris*.

Figure 7 provides a representation of pHSA113 in linear form.

Figure 8 provides a representation of plasmid pA0807N which contains a linear site-specific integrative vector in the fragment clockwise from Notl to Notl. The structural gene may be inserted in the unique EcoRl site of this plasmid.

35 Detailed Description

The present invention provides improved expression cassettes for the expression of HSA, improved vectors and *Pichia pastoris* strains transformed with these improved cassettes and vectors.

Utilizing the present invention, HSA secretion levels of approximately 1-3.4 grams of authentic HSA per to liter of fermentation broth have been obtained with an additional 15-88 µg HSA per mg of protein being present within the cell secretory pathway. This invention thus provides a means for the high level secretion of HSA. Achieving these levels of HSA production is a significant advancement over the prior production levels, since at the level of 1-3.4 grams per liter the recovery of HSA in high yields with high purities is possible.

To express the HSA structural gene, the gene must be operably linked to a 5' regulatory region and a 3' termination sequence, which forms an expression cassette which will be inserted into a host (usually a microorganism) via a vector (such as a circular plasmid or linear site-specific integrative vector). Operably linked as used in this context refers to a juxtaposition wherein the 5'regulatory region, structural gene, and 3' termination sequence are linked and configured so as to perform their normal function. 5' regulatory region or promoter as used herein means DNA sequences which respond to various stimuli and provide enhanced rates of mRNA transcription. 3' termination sequences are equences 3' to the stop codon of a structural gene which function to stabilize the mRNA transcription product of the gene to which this invention, it is preferred that the ATG of the structural gene be linked with as few intervening deox-structured production of the structural gene be linked with as few intervening deox-structured production of the structural gene be linked with as few intervening deox-structured production of the structural gene be linked with as few intervening deox-structured production of the structural gene be linked by the structural gene when the structural gene which function to the structural gene be linked with as few intervening deox-structured gene which functions the structural gene be linked by the structural gene be linked to the structural gene

locations. Counting lett from the ATG codon of the structural gene with the first position let being the -1 position it appears that adenine or cytosine is the most preferred deoxyribonucleotide, in the -2 position the most preferred deoxyribonucleotide is either adenine or thymine, in the -3 position the most preferred deoxyribonucleotide is adenine or thymine and the most preferred uncleotide at the -4 position is adenine, s thymine or cytosine. Currently, it is preferred that the ADX or DAS 15' regulatory regions having the restriction maps of Figures 3 and 4 or, the sequences provided as SEQ ID No: 1 and SEQ ID No: 2, respectively, be linked at their 3' end of the sequence to the ATG start codon of the HSA structural gene. Two examples of appropriate linkages for the ADX15' resulatory region are illustrated below.

1	Construct Designation	End of the 5' Regulatory Region for AOX 1	Deoxyribonucleotide intervening before ATG start condon
	pHSA140	5' - TTCGAAACG	5' - AGGAATTC
	pHSA413, pHSA313	5' - TTCGAAACG	5' - NONE

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Several regulatory regions have been characterized and can be employed in conjunction with the expression of HSA in *Pichia pastoris*. Exemplary 5' regulatory regions are the primary alcohol oxidase (AOX1), dihydroxyacetone synthase (DAS1), and the p40 regulatory regions, derived from *Pichia pastoris* and the like. The presently preferred 5' regulatory regions employed in the practice of this invention are those characterized by their ability to respond to methanol-containing media, such regulatory regions selected from the group consisting of AOX1, and DAS1, disclosed by D. W. Stroman et al. in U.S. Patent 4,855,231, incorporated herein by reference. The most preferred 5' regulatory region for the practice of this invention is the AOX1 5' regulatory region.

3' termination sequences should be utilized in the expression cassette as discussed above. 3' termination sequences smy function to terminate, polyadenylate and/or stabilize the messenger RNA coded for by the structural gene when operably linked to a gene, but the particular 3' termination sequence is not believed to be critical to the practice of the present invention. A few examples of illustrative sources for 3' termination sequences for the practice of this invention include but are not limited to the Hansenular polymorpha and Pichia pastoris 3' termination sequences. Preferred are those derived from Pichia pastoris such as those selected from the group consisting of the 3' termination sequence of ADX1 gene.

DAS1 gene, p40 gene and HIS4 gene. Particularly preferred is the 3' termination sequence of the ADX1.

Phila pastoris may be transformed with a variety of HSA structural genes (in the inventive transformants discussed herein the HSA structural gene encodes both a signal sequence and a mature HSA protein). HSA structural genes have been sequenced by Lawn et al. Nuc. Acids Res. 9:6105 (1981), and Dugaiczyk et al., Proc. Natl. Acad. Sci. USA 79:71 (1982). These genes may also be obtained by reisolation of the genes by the Technique of Lawn et al., Dugaiczyk et al. or synthesized in vitro by a custom gene manufacturer such as British Biotechnology, Ltd. One possible method of obtaining \$\overline{a}\$ HSA gene would be to screen a human liver cDNA interry with oligonucleotide probes or screen a human liver cDNA interry with oligonucleotide probes or screen a human liver cDNA interry with oligonucleotide probes or screen a human liver cDNA interry with oligonucleotide probes or screen a human liver cDNA interry with oligonucleotide probes or screen a human liver cDNA interry with oligonucleotide probes or screen a human liver cDNA interry with oligonucleotide probes or screen a human liver cDNA interry with oligonucleotide probes or screen a human liver cDNA interry with oligonucleotide probes or screen a human liver cDNA interry with oligonucleotide probes or screen a human liver cDNA interry with oligonucleotide probes or screen a human liver cDNA interry with oligonucleotide probes or screen a human liver cDNA interry with oligonucleotide probes or screen a human liver cDNA interry with oligonucleotide probes or screen a human liver cDNA interret value of the probes of the pr

Plasmid type vectors have long been one of the basic elements employed in recombinant DNA technology. Plasmids are circular extra-chromosomal double-stranded DNA found in microorganisms. Plasmids have been found to occur in single or multiple copies per cell. Included in plasmid DNA is the information required for plasmid reproduction, e.g. an autonomous replication sequence such as those disclosed by James M. Cregg in U.S. Patent 4,837,148, issued June 6, 1989, incorporated herein by reference. The autonomous replication sequences disclosed by Cregg provide a suitable means for maintaining plasmids in Pichia pastoris. Additionally one or more means of phenotypically selecting the plasmid in the plasmid.

Suitable integrative vectors for the practice of the present invention are the linear site-specific integrative vectors described by James M. Cregg, in U.S. Patent 4,882.279, issued November 21, 1889, which is incorporated herein by reference. These vectors comprise a serially arranged sequence of at least 1) a first insertable DNA fragment; 2) a selectable marker gene; and 3) a second insertable DNA fragment. An expression cassette containing a heterologous structural gene is inserted in this vector between the first and second insertable DNA fragments either before or after the marker gene. Alternatively, an expression cassette can be formed in 81 fix if a regulatory region or promoter is contained within one of the insertable

fragments to which the structural gene may be operably linked.

The first and second insertable DNA fragments are each at least about 200 nucleotides in length and have nucleotide sequences which are homologous to portions of the genomic DNA of the species to be transformed. The various components of the integrative vector are serially arranged forming a linear fragment of DNA such that the expression cassette and the selectable marker gene are positioned between the 3' end of the first insertable DNA fragment and the 5' end of the second insertable DNA fragment. The first and second insertable DNA fragments are oriented with respect to one another in the serially arranged linear fragment as they are oriented in the parent genome.

Nucleotide sequences useful as the first and second insertable DNA fragments are nucleotide sequences which are homologous with separate portions of the native genomic site at which genomic modification is to occur. For example, if genomic modification is to occur at the locus of the alcohol oxidase gene, the first and second insertable DNA fragments employed would be homologous to separate portions of the alcohol oxidase gene locus. Examples of nucleotide sequences which could be used as first and second insertable DNA fragments are deoxyribonucleotide sequences selected from the group consisting of 15 the Pichia pastoris alcohol oxidase (AOX1) gene, dihydroxyacetone synthase (DAS1) gene, p40 gene and HIS4 gene. The AOX1 gene, DAS1 gene, p40 gene and HIS4 genes are disclosed in U.S. Patents 4,855,231 and 4,885,242 both incorporated herein by reference. The designation DAS1 is equivalent to the DAS designation originally used in U.S. Patents 4,855,231 and 4,885,242.

The first insertable DNA fragment may contain an operable regulatory region which may comprise the regulatory region utilized in the expression cassette. The use of the first insertable DNA fragment as the regulatory region for an expression cassette is a preferred embodiment of this invention. Figure 1 provides a diagram of a vector utilizing the first insertable DNA fragment as a regulatory region for a cassette. Optionally, as shown in Figure 1, an insertion site or sites and a 3' termination sequence may be placed immediately 3' to the first insertable DNA fragment. This conformation of the linear site-specific integrative 25 vector has the additional advantage of providing a ready site for insertion of a structural gene without necessitating the separate addition of a compatible 3' termination sequence.

If the first insertable DNA fragment does not contain a regulatory region, a suitable regulatory region will need to be inserted linked to the structural gene, in order to provide an operable expression cassette. Similarly, if no 3' termination sequence is provided at the insertion site to complete the expression cassette, a 3' termination sequence can be operably linked to the 3' end of the structural gene.

It is also highly desirable to include at least one selectable marker gene in the DNA used to transform the host strain. This facilitates selection and isolation of those organisms which have incorporated the transforming DNA. The marker gene confers a phenotypic trait to the transformed organism which the host did not have, e.g., restoration of the ability to produce a specific amino acid where the untransformed host strain has a defect in the specific amino acid biosynthetic pathway, or provides resistance to antibiotics and the like. Exemplary selectable marker genes may be selected from the group consisting of the HIS4 gene (disclosed in U.S. Patent 4,885,242) and the ARG4 gene (disclosed in U.S. Patent 4,818,700 incorporated herein by reference) from Pichia pastoris and Saccharomyces cerevisiae, the invertase gene (SUC2) (disclosed in U.S. Patent 4,857,467 incorporated herein by reference) from Saccharomyces cerevisiae, or the G418^R/kanamycin resistance gene from the E. coli transposable elements Tn601 or Tn903.

Those skilled in the art recognize that additional DNA sequences can also be incorporated into the vectors employed in the practice of the present invention, such as, for example, bacterial plasmid DNA, bacteriophage DNA, and the like. Such sequences enable the amplification and maintenance of these vectors in bacterial hosts.

The insertion of the HSA structural gene into suitable vectors may be accomplished by any suitable technique which cleaves the chosen vector at an appropriate site or sites and results in at least one operable expression cassette containing the HSA structural gene being present in the vector. Ligation of the HSA structural gene may be accomplished by any appropriate ligation technique such as utilizing T4 DNA

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The initial selection, propagation, and optional amplification of the ligation mixture of the HSA structural gene and a vector is preferably performed by transforming the mixture into a bacterial host such as E. coli -(although the ligation mixture could be transformed directly into a yeast host but, the transformation rate would be extremely low). Suitable transformation techniques for E. coli are well known in the art. Additionally, selection markers and bacterial origins of replication necessary for the maintenance of a vector 55 in a bacterial host are also well known in the art. The isolation and/or purification of the desired plasmid containing the HSA structural gene in an expression system may be accomplished by any suitable means for the separation of plasmid DNA from the host DNA. Similarly the vectors formed by ligation may be tested, preferably after propagation, to verify the presence of the HSA gene and its operable linkage to a regulatory region and a 3' termination sequence. This may be accomplished by a variety of techniques including but not limited to endonuclease digestion, gel electrophoresis, or Southern hybridization.

Transformation of plasmids or linear vectors into yeast hosts may be accomplished by suitable transformation techniques including but not limited to those taught by Cregg and Barringer, U.S. Patent 4,929,555; Hinnen et al., Proc. Natl. Acad. Sci. 75, (1978) 1929; Ito et al., J. Bacteriol. 153, (1983) 183; Cregg et al. Mol. Cell Biol. 5 (1985), pp. 3376; D. W. Stroman et al., U.S. Patent 4,679,231, issued November 7, 1989; or Sirevishna et al., ene, 59 (1987), pp. 115. Preferable for the practice of this invention is the transformation technique of Cregg et al. (1985). It is desirable for the practice of this invention is utilize linear vectors and select for insentions by Southern hybridization.

The yeast host for transformation may be any suitable methylotrophic yeast. Suitable methylotrophic yeasts include but are not limited to yeast capable of growth on methanol selected from the group consisting of the genera *Hansenula* and *Pichia*. A list of specific species which are evemplary of this class of yeasts may be found in C. Anthony, The Biochemistry of Methylotrophs, 289 (1982). Presently preferred are methylotrophic yeasts of the genus *Pichia* past for Bichia past for SG115 (NRRL Y-18014) disclosed in U.S. Patent 4,812,700; and *Pichia* past for SF15); *Pichia* past for SG150 (NRRL Y-18014) disclosed in U.S. Patent 4,812,700; and *Pichia* past for SF15); *Pichia* past for SF15 (NRRL Y-18014) disclosed in U.S. Patent 4,812,700; and *Pichia* past for SF15); *Pichia* past for SF15 (NRRL Y-18014) disclosed in U.S. Patent 4,812,700; and *Pichia* past for SF15); *Pichia* past for SF15 (NRRL Y-18014) disclosed in U.S. Patent 4,812,700; and *Pichia* past for SF15); *Pichia* past for SF15 (NRRL Y-18014) disclosed in U.S. Patent 4,812,700; and *Pichia* past for SF15); *Pichia* past for SF15 (NRRL Y-18014) disclosed in U.S. Patent 4,812,700; and *Pichia* past for SF15); *Pichia* past for SF15 (NRRL Y-18014) disclosed in U.S. Patent 4,812,700; and *Pichia* past for SF15); *Pichia* past for SF15 (NRRL Y-18014); *Pichia*

Transformed Pichia pastoris cells can be selected for by using appropriate techniques including but not limited to culturing previously auxotrophic cells after transformation in the absence of the biochemical product required (due to the cell's auxotrophy), selection for and detection of a new phenotype ("methanol slow"), or culturing in the presence of an antibiotic which is toxic to the yeast in the absence of a resistance gene contained in the transformant.

Isolated transformed *Pichia pastoris* cells are cultured by appropriate fermentation techniques such as shake flask fermentation, high density fermentation or the technique disclosed by Crege et al. in, <u>Hight-velov</u> Expression and Efficient Assembly of Hepatitis B Surface Antigen in the Methylotrophic Yeast, *Pichia Pastoris* 5 Bio/Technology 479 (1987), Isolates may be screened by assaying for HSA production to identify those isolates with the highest HSA production level.

Transformed strains, which are of the desired phenotype and genotype, are grown in ferrmentors. For the large-scale production of recombinant DNA-based products in methylotrophic yeast, a three stage, high cell-density, batch fermentation system is normally the preferred fermentation protocol employed. In the first, or growth stage, expression hosts are cultured in defined minimal medium with an excess of a non-inducing carbon source (e.g. glycerol). When grown on such carbon sources, heterologous gene expression is completely repressed, which allows the generation of cell mass in the absence of heterologous protocomic expression. It is presently preferred, during this growth stage, that the pH of the medium be maintained at about 5. Next, a short period of non-inducing carbon source limitation growth is allowed to further increase cell mass and deerpress the methanol responsive promoter. The pH of the medium during this limitation growth at about pH 5.6 to about pH 6.5, referably either about pH 5.0 or about pH 5.8. Subsequent to the period of growth under limiting conditions, methanol alone (referred to herein as "limited methanol fed-batch-mode") or a limiting amount of non-inducing carbon source plus methanol (referred herein as "limited methanol fed-batch-mode") are added in the fermentor, inducing the expression of the heterologous gene driven by a methanol responsive promoter. This third stage is the so-called production stage.

The invention will now be described in greater detail in the following non-limiting examples.

Examples

General information pertinent to the Examples:

Strains

Pichia pastoris GS115 (his 4) NRRL Y-15851

E, coli JM103 delta (lac pro) thi rpsl (strA) supE endA sbcB hsdR.

E. coli K12 MC1061 NRR1-18016 (F⁻, araD139 delta (lac 1POZY)x74 galk galu hsr hsm(+) rpsL delta (araABOIC leu)7697.

E. coli DG75' (hsd1, leu6, lacY, thr-1, supE44, tonA21, lambda-)

Buffers, Solutions and Media

5	The buffers, solutions, and media employed in the below:	e following examples have the compositions given
	dH₂O	deionized H ₂ O that has been treated with a milli-Q (Millipore) reagent water system.
	and Trie hulles	121.1 g Tris base in 800 mL of H ₂ O; adjust
	1M Tris buffer .	pH to the desired value by adding concen-
10		trated (35%) aqueous HCl; allow solution to
		cool to room temperature before final pH ad-
		justment, dilute to a final volume of 1 L.
	TE buffer	1.0 mM EDTA
15	12 001101	in 0.01 M (pH 8.0) Tris bufffer
,,	SED	1 M sorbitol
	020	25 mM EDTA
		50 mM DTT, added prior to use
		adjust to pH 8
20	SCE	9.1 g sorbitol
		1.47 g Sodium citrate
		0.168 g EDTA
		pH to 5.8 with HCl in 50 ml
		dH₂O and autoclave
25	CaS	1 M sorbitol
		10 mM CaCl ₂
		filter sterilize
	SOS:	1 M sorbitol 0.3x YPD
		10 mM CaCl ₂
30	000	20% polyethylene glycol-3350
	PEG	10 mM CaCl ₂
		10 mM Tris-HCl (pH 7.4)
		filter sterilize
35	Solution A	0.2 M Tris-HCl (pH 7.5)
33	Soldion A	0.1 M MgCl ₂
		0.5 M NaCl
		0.01 M dithiothreitol (DTT)
	Solution B	0.2 M Tris-HCI (pH 7.5)
40		0.1 M MgCl ₂
		0.1 M DTT
	Solution C (keep on ice)	4 μl solution B
		4 μl 10 mM dATP
		4 μl 10 mM dTTP
45		4 μl 10 mM dGTP
		4 µl 10 mM dCTP
		4 µl 10 mM ATP
		5 µl Т₄ ligase (2 U/µl) 12 µl H₂O
		Recipe for Solution C was modified from Zol-
50		ler & Smith
	201/ 2005	4.4 g NaOH
	20X SSPE	7.4 g Na ₂ EDTA
		27.6 g NaH ₂ PO ₄ *H ₂ O
		210 g NaCl
55		pH adjusted to 7.5-8.0 with NaOH
		H ₂ O to 1 liter
	50X Denhardt's	5 a Ficoli 400
		•

		5 g Polyvinylpyrolidine
		5 g BSA Fraction V
		H ₂ O to 500 ml
	20X SSC	175.3 g NaCl
		88.2 g sodium citrate
		pH to 7.0 with NaOH
		H ₂ O to 1 liter
	LB Broth, 1 liter	5.0 g yeast extract
	·	10.0 g tryptone
		5.0 g NaCl
	10X Transfer Buffer	96.8 g Trizma Base
	TOX TRANSPORTE	9.74 g glycine
		water to 1 liter
	Transfer Buffer for Tank	500 mls 10X Transfer Buffer
	Transier Boner for Tank	1000 mls methanol
		3500 mls water
	Western Buffer - for 1 liter	2.5 g gelatin put in solution by microwaving
	Western Dulier - IOI 1 lifel	first in 100 mls water
	•	100 mls 10X PBS
		1 ml 50% Tween-20
	·	4 mls 5% sodium azide
	0 4 9 4	dH ₂ O to 1 liter
	Coating Buffer	0.160 g Na ₂ CO3 (sodium carbonate)
		0.294 g NaHCO3 (sodium carbonate)
		Add distilled water to 100 ml. Do not pH. (pH
		should be 9.5)
	Tris Buffered Saline (TBS)	26.1 g NaCl
		2.63 g Tris
		Add distilled water to 3 liters.
•		Adjust pH to 7.5 with HCl.
	Tris Buffered Saline/Tween (TBST)	1 liter of TBS
		2.5 ml of 20% Tween-20
	Blotto Buffer	50 g of non-fat dry milk (Carnation)
		1 g thimerosal (Sigma)
	7 1	100 µl of antifoam (Sigma, 30% emulsion)
		2.5 ml of 20% Tween-20
		100 ml 10x PBS (house stock)
		Add distilled water to 1 liter
		Adjust pH to 7.5
)	Ligation Buffer	50 mH Tris-HCl (pH 7.4)
		10 mM MgCl ₂
		10 mM dithiothreitol
		1 mM ATP
	Phosphatase Buffer	50 mM Tris-HCl (pH 9.0)
5		1 mM MgCl ₂
		1 mM ZnCl ₂
		1 mM spermidine
	Bsu36l buffer	100 mM NaCl
		10 mM Tris-HCl (pH 7.4)
)		10 mM MgCl ₂
		100 µg/ml BSA
	Csp45I buffer	60 mM NaCl
		10 mM Tris-HCl, pH 7.5
		7 mM MgCl ₂
5		100 μg/ml BSA
	REact 1 buffer	50 mM Tris-HCl, pH 8.0
	- 12	10 mM MgCl ₂
		100 μg/ml BSA
		-

	REact 2 buffer	REact 1 buffer + 50 mM NaCl
	REact 3 buffer	REact 1 buffer + 100 mM NaCl
	HS buffer	50 mM Tris-HCl, pH 7.5
		10 mM MgCl ₂
5		100 mM NaCl
		1 mM DTT
		100 µg/ml BSA
	10X Basal Salts	42 mls Phosphoric Acid, 85%
		1.8 g Calcium Sulfate * 2H ₂ O
10		28.6 g Potassium Sulfate
		23.4 g Magnesium Sulfate * 7H ₂ O
		6.5 g Potassium Hydroxide
	Ptm ₁ Trace Salts Solution	C.O Cunda Sulfata + EM.O
		6.0 g Cupric Sulfate * 5H ₂ O
15		0.08 g Sodium lodide
		3.0 g Manganese Sulfate * H ₂ O
		0.2 g Sodium Molybdate * H ₂ O 0.02 g Boric Acid
		0.5 g Cobalt Chloride
		20.0 g Zinc Chloride
20		65.0 g Ferrous Sulfate * H ₂ O
		0.20 g Biotin
		5.0 mls Sulfuric Acid
	YPD (yeast extract peptone dextrose medium)	3.0 This Culture Acid
25	TPD (yeast extract peptone dextrose mediani)	10 g bacto yeast extract
20		20 g peptone
		10 g dextrose
		water to 1 liter
	MGY (minimal glycerol medium)	
30	,	13.4 g yeast nitrogen base with ammonium
		sulfate, and without amino acids
		400 μg biotin
		10 ml glycerol
		water to 1 liter
35	MM (minimal methanol medium)	•
		Same as MGY, except that 5 ml methanol is
		used in the place of 10 ml glycerol.
	SDR (supplemented dextrose regeneration medium):	
		13.4 g yeast nitrogen base with ammonium
40		sulfate and without amino acids
		400 μg biotin
		182 g sorbitol
		10 g glucose
		2 g Histidine assay mix (Gibco)
45		50 mg glutamine
		50 mg methionine
	8 .	50 mg lysine
		50 mg leucine
		50 mg isoleucine
50		10 g agarose
	DAKED (D. Hand minimal almost a parished medium)	water to 1 liter
	BMGR (Buffered minimal glycerol-enriched medium)	100 ml/liter Potassium phosphate buffer, (pH
		6.0)
		13.4 grams/liter Yeast nitrogen base with am-
55		monium sulfate
		400 µg/liter biotin
		10 ml/liter glycerol
		gijosio.

Amino acids

glutamic acid, methionine, lysine, leucine and isoleucine: each at 5 mg/liter;

all the other amino acids except histidine at 1 mg/liter

Nucleotides Vitamins

adenine sulfate, guanine hydrochloride, uracil, and xanthine, each at 40 µg/liter

thiamine hydrochloride, riboflavin, and calcium

pantothenate, each at 2 µg/liter; pyridoxide hydrochloride and nicotinic acid,

each at 4 µo/liter: pyridoxamine hydrochloride and pyridoxal hydrochloride, each at 1 µg/liter;

para-amino benzoic acid at 0.3 µg/liter;

folic acid at 0.03 µg/liter

Trace minerals

magnesium sulfate at 800 µg/liter; ferrous sulfate at 40 µg/liter; manganese sulfate at 80 µg/liter; sodium chloride at 40 µg/liter

BMGY (Buffered minimal glycerol-complex medium)

100 ml/liter potassium phosphate buffer, (pH 6.0)

13.4 grams/liter yeast nitrogen base with ammonium sulfate and without amino acids biotin at 400 µg/liter glycerol at 10 ml/liter

veast extract at 10 g/liter peptone at 20 g/liter

BMMR (Buffered minimal methanol-enriched medium)

Same as BMGR, with the exception that 5 ml methanol/liter is added in the place of glycerol Same as BMGY, with the exception that 5 ml methanol/liter is added in the place of glycerol

BMMY (Buffered minimal methanol -complex medium)

Techniques

35

Suitable techniques for recombinant DNA lab work may be found in many different references including but not limited to: Methods in Enzymology, (Orlando, FL: Academic Press, Inc.), particularly Volume 152, published as, Guide to Molecular Cloning Techniques, by Berger and Kimmel (Orlando, FL: Academic Press, Inc., 1987) and Molecular Cloning/A Laboratory Manual, by Sambrook et al., 2d ed. (Cold Spring Harbor Laboratory Press, 1989) and which are all hereby incorporated by reference.

Example 1

Construction of mHSA13

Mutagenesis of HSA Structural Gene Insert

DNA encoding HSA was obtained from pHSA13, disclosed in European Patent Application 0 344 459, herein incorporated by reference, by EcoRI digestion. A 2069 bp fragment was recovered by electrophoresis on a 1% agarose gel. The DNA was mutagenized by the following procedure to make the following changes: 1) an EcoRI restriction site was added immediately prior to the ATG of the HSA signal sequence, and 2) an EcoRI restriction site was added immediately adjacent to the TAA stop codon in the HSA cDNA.

The oligonucleotides employed in the mutagenesis were:

1) 5' mutagenesis to add EcoRI site, mutagenizing nucleotide sequence:

5' CCC TCA CAC GCC TTT GAA TTC ATG AAG TGG GTA ACC 3' (SEQ ID NO:4)

2) 3' mutagenesis to add EcoRI site, mutagenizing nucleotide sequence:

5' GCC TTA GGC TTA TAA GAA TTC AGT TTA AAA GCA TCT CAG 3' (SEQ ID NO:5)

and were synthesized using an Applied Biosystems DNA Synthesizer, Model 380A using cyanoethyl-phosphoramidite chemistry.

 $1.2~\mu g$ of double-stranded m13mp10 were digested with EcoRI and dephosphorylated and ligated with 450 ng of the previously isolated 2069 bp fragment containing the HSA structural gene.

The ligation mixture was transformed into competent JM103 cells (competent JM103 were prepared as described in Example II for MC1061 cells). The mixture was then plated on LB media containing IPTG and X-gal and the plates screened for clear plaques. DNA was recovered from transformants and digested with Hind III. The correct phage demonstrated bands of 7369 and 1950 bp and was called mHSA13.

A. A large scale miniprep was performed on positive plaques which had been incubated for approximately 7 hours in 2 mils of 1. media. 25 mils of LB media was incoulated with 250 µl of freshly grown JM103 cells. The culture was grown for 1 hour and incoulated with 100 µl of the 7 hour old plaque culture. The culture was then grown overnight. The culture was centrifuged twice at 10,000 rpms for 10 minutes on a Soval RC-SB rotor SS34 to clear the supernation 3.5 mil of 20% PEGI2.5 M NaCl was added to the culture and it was incubated for 5 hours at 4 °C. The culture was then centrifuged again as above for 10 minutes. The supernatant was discarded and the pellet was resuspended in 2 mils of TE buffer. The pellet was suspended in 2 mils of TE buffer. The pellet was resuspended in 2 mils of TE with a continuation of 10.8 M. 3 volumes of ethanol were added and the solution left overnight at 20°C to precipitate the DNA present. The solution was next centifuged for 10,000 rpms for 10 minutes as previously described and rinsed with 75% ethanol. The precipitate was resuspended in 150 µl of 10 mM Tris (pH 7.4).

B. One pmole of M13 recombinant template was mixed with 20 pmole of oligonucleotide 1 (for 5' mutagenesis to create an EcoRl site), 1 μl of solution A and dH₂O was added to give a final volume of 10 μl. The sample was incubated at 65° C for 5 minutes, and the temperature was then reduced to 37° C for 30 minutes.

C. The following was then added to the sample:

Solution B	1 μΙ
10 mM dATP	1 µl
10 mM dCTP	1 µl
10 mM dGTP	1 µl
10 mM dTTP	1 µl
5 u/μl Klenow	2 µl
dH₂O	3 41
	20 µI

and allowed to incubate at 15 °C for at least 4-6 hours.

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D. The sample was then diluted 1:40 with dH_2O , 5 μl was used to transform 6 tubes of competent JM103 cells (200 μl each). The transformed JM103 cells were plated on rich media in a soft agar overlay.

E. The positive plaques were then screened for by filter hybridization.

A hybridization probe of 15 pmole of complementary oligonucleotide in a total volume of 25 μl total volume was heated to 65°C for 10 minutes. 3 μl 10X kinase buffer (Maniatis), 1 μl γ-ATP and 1 μl polynucleotide kinase (100 μ/μl) were added to the sample. The sample was incubated for 1 hour at 37°C and run through 6-50 fine Sephadex. The first peak off the column was collected.

Nitrocellulose filters were prepared for hybridization with the above probe by placing and orienting the filters on the transformation plates for 5-10 minutes. The filters were then removed from the plates and floated on a denaturing solution (1.5 M NaCl, 0.5 N NaCl) for 3 minutes with the backside on top of the solution. The filters were then submerged in the denaturing solution for 5 minutes. The nitrocellulose filters were transferred to a neutralizing solution (1 M Tris' H/Cl, pH 8; 1.5 M NaCl) for 5 minutes. The neutralized filter was then transferred to 2XSSC (1XSSC is 150 mM NaCl, 15 mM NaClitrate) for 5 minutes. The filter was then transferred to 2XSSC (1XSSC is 150 mM NaCl, 15 mM NaClitrate) for 5 minutes. The filter was then air dried and baked for 1 hour at 80 °C under a vacuum. The filters were prehybridized for 1 hour at 65 °C in a sealed plastic bag containing 5 ml of hybridization buffer filter, 10X Denhardts (1X Denhardts is 0.02% Ficoll, 0.02% polyvinyl pyrrolidone, 0.02% bovine serum albumin) 0.5% SDS and SXSSPE. The hybridization buffer was replaced with 5 miltitler of fresh hybridization buffer was replaced with 5 miltitler of fresh hybridization buffer was replaced with 5 miltitler of fresh hybridization buffer was replaced with 5 miltitler of fresh hybridization buffer was replaced with 5 miltitler of fresh hybridization buffer was replaced with 5 miltitler of fresh hybridization buffer was replaced with 5 miltitler of fresh hybridization buffer was replaced with 5 miltitler of fresh hybridization buffer was replaced with 5 miltitler of fresh hybridization buffer was replaced with 5 miltitler of fresh hybridization buffer was replaced with 5 miltitler of fresh hybridization hybridization buffer was replaced with 5 miltitler of fresh hybridization hybridiza

buffer. The screening oligonucleotides utilized were as follows:

- 1) 5' mutagenesis the screening oligonucleotide was
- 5' GCC TGG GAA TTC ATG AAG 3' (SEQ ID NO:6)
- 2) 3' mutagenesis the screening oligonucleotide was 5' TTA TAA GAA TTC AGT TTA 3' (SEQ ID NO:7)

The previously prepared screening oligonucleotide was first incubated at 65°C for 5 minutes, and then enough probe was added to the fresh hybridization buffer containing the filter to give 1X10⁶ cpm/ml. Hybridization was performed at 5°C below the calculated melting temperature of the probe for 4 hours.

The filters were then washed three times for 10 minutes each with 6XSSC at room temperature. The filters were finally asshed one time with 6XSSC at the hybridization temperature. The filters were placed on a 3 MM Whatman paper to dry, and then exposed to film (marked for orientation) overnight.

Three positive plaques were each picked and grown separately in 2 mls of LB broth at 37 °C for 5

F. Mini template preps were performed on each of these positive plaques.

One mI of the pisque culture was transferred into an Eppendorf tube and centrifuged for 5 minutes in a Eppendorf Model S414 Centrifuge. 800 µl of the supernatant was recovered and 200 µl of 20% PEG with 2.5M NaCl was added thereto. The supernatant was incubated at room temperature for 10 minutes. The supernatant was centrifuged for 10 minutes in the Eppendorf centrifuge previously used. The supernatant was removed by aspiration and the pellet formed by centrifuging was redissolved in 200 µl TE (10 mM Tris, pH 7.4; 1 mM EDTA). The redissolved pellet was then phenol/chloroform extracted and the template DNA in the upper aqueous phase was precipitated by the addition of a LICI solution until a 0.8 M concentration was reached. To the solution was added 2 172-3 volumes of ethanol and precipitated on dry ice for 5 minutes. The precipitate was centrifuged for 10 minutes in the previously mentioned Eppendorf centrifuce. The final volume was brought up to 150 µl with TE.

G. 200 μ I of competent JM103 cells were transformed with the recovered DNA. 1 μ I and 1 μ I of a 1/10 dilution of the isolated phase DNA was used in the transformation.

H. The transformation mixture was plated and plaques were screened with oligonucleotides as previously described in step E.

I. A large scale miniprop was performed on positive plaques which had been incubated for approximately 7 hours in 2 mls of L media. 25 mls of LB media was inoculated with 250 µl of freshly grown JM103 cells. The culture was grown for 1 hour and inoculated with 100 µl of the 7 hour old plaque culture. The culture was then grown overnight. The culture was then centrifuged twice at 10.000 prms for 10 minutes on a Sorvall Rf-C5 protor S334 to clear the supernatant. 35 ml of 20% PEG/2.5M NaCl was added to the culture and it was incubated for 5 hours at 4 °C. The culture was then centrifuged again as above for 10 minutes. The supernatant was discarded and the pellet was resuspended in 2 mls of TE buffer. The pellet was then extracted with phenol/chloroform extracted brice with CHCl₃ and once with ether. 8 M LICI was added to attain a final concentration of 0.8 M LiCl. 3 volumes of ethanol were added and the solution left overright to precipitate the DNA present. The solution was next centrifuged for 10,000 rpms for 10 minutes as proviously described and rinsed with 70% ethanol. The precipitate was resuspended in 150 µl of 10mM Tris (eH7-4).

J. The positive plaques were then sequenced by dideoxy sequencing to find the M13 constructs with the correct mutations.

K. Repeat steps B-J using M13 constructs with the correct 5' mutations as templates and the second oligonucleotide as a primer for 3' mutagenesis. The correct mutation was designated mHSA140.

L. Recover RF DNA of mHSA140 using the alkaline lysis method of Maniatis.

Example II

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Construction of the pHSA140 Expression Vectors

pA0804 is available in an *E. coli* host from the Northern Regional Research Center of the United States Department of Agriculture, Peoria, Illinois, accession number B-18114. pA0804 is recovered by isolating the plasmid DNA, digesting with EcoRI, gel electrophoresing to recover the ~7.5 kb fragment, which is linear pA0804 cut at its unique EcoRI site.

pA0804 is a vector capable of site-specific disruption of the *Pichia pastoris* <u>AOX1</u> locus. It contains the following elements: the AOX1 promoter and transcription terminator separated $\frac{1}{2}$ unique EcoRl cloning site; the wild-type *Pichia* HiS4 gene; a genomic segment of DNA from the 3' end of the AOX1 locus downstream of the transcription terminator; and sequences necessary for selection and replication in a

bacterial host. The components are arranged such that a BgIII restriction digest of the plasmid releases a DNA fragment containing the expression cassette and selective marker whose ends are homologous to a continuous portion of the genome, the AOXI locus, and can be stably inserted into the chromosome during transformation. Additionally the ampicillin resistance gene and the orl from plasmid pBR322 are also contained in the pA0804 plasmid.

A vector containing the gene coding for the production of HSA was constructed from pA0804 and mHSA140. pA0804 was digested with EcoRI and the ends were dephosphore/lated by treatment with alkaline phosphatase (1 U enzyme at 37° C for T in: 50 mM Tris* CI, pH 9.0, 1 mM MgOz, 100 mM ZnClp, 1 mM spermidine), mHSA140 was also digested with EcoRI, and a 1829 by fragment encoding HSA was released. This fragment was purified using 0.8% preparative agarose gel electrophoresis. 60 ng of the fragment were ligated to 240 ng of pA0804 by incubation at 23° C for 1 hr in 66 mM Tris* CI, pH 7.4, 5 mM MgClp, 5 mM dithiotheriol, 1 mM ATP, with 1 Weiss Unit of 14 ligase in a 10 µI reaction volume. The ligation reaction was used to transform competent MC1061 cells to amplicilin resistance.

MC1061 was rendered competent for transformation in the following manner. A mid-log culture (50 ml)
s of *E. coli* MC1061 was harvested by centrifugation in an IEC DPR 600 clinical centrifuge at 3,000 rpm for 5 min at 4 °C and washed in 10 mM NaCl. The culture was resuspended in 25 ml of 50 mM CaCl₂ for 30 min at 0 °C. The cells were centrifuged as above and resuspended in 2 ml of 50 mM CaCl₂.

For transformation, the ligation reaction was added to 200 µl of the competent cell suspension and incubated at 0 °C on ice for 15 minutes, heat shocked at 37 °C for 5 minutes and incubated at 23 °C for 5 and provided to 123 °C for 5 minutes and incubated at 23 °C for 50 minutes. The cells were plated directly onto LB agar plates containing 50 µg/ml ampicillin. The plates were incubated at 37 °C for 10-16 hours. The resulting colonies were Ampl⁸. The resistant colonies were harvested and characterized by restriction digestion. Cells were grown in 5 ml of 1-borth containing 50 µg/ml ampicillin for 5 hr at 37 °C and DNA was prepared by the method of Birnboim and Doly [Nucleic Acids Research 7:1513 (1979)]. The minipreps displaying 4750, 3000 and 1900 bp fragments upon Pvull 25 digestion were chosen and designated PtSN140.

Example III

Transformation of Pichia pastoris

Pichia pastoris strains containing the vectors described in Example II were generated in the following manner. Methanol utilization deficient (Mut* or methanol slow) and wild type methanol utilization (Mut* or methanol ormal) strains were developed.

Pichia pastoris strain GS115 (his4; NRRL Y-15851) was transformed using the spheroplast transformass tion technique described by Cregg et al., Bio/Technology 5;479-485 (1987). See also U.S. 4,879,231.

A. Mut Strains

To direct integration of the vector to the AOX1 locus, 2 and 10 μg of Sac1- digested pHSA140 were 40 separately transformed into 5 of Obco (or 25 x 10 cells) of GS115. Transformants were regenerated on minimal media and screened for the His phenotype. Several His transformants were then screened by Southern analysis for the site of integration and vector copy number (Example V).

B. Mut Strains

To develop Mut" strains, in which the HSA expression cassette inegrates into and disrupts the AOX1 structural gene, vector pHSA140 was digested with Pvul and then partially digested with Bglll. The digest was then size fractionated on a 0.8% agazose gel and DNA in the size range of 6.0-9.0 kb was isolated (the expression cassette was expected to be " 7.4 kb). 5µg of this DNA were used to transform 5 ODeco (25 x 50 10) cells) of GS115 by the spheroplast method. His cells were identified and then screened for the Mut" phenotive as follows.

Transformants were pooled by scraping the surface of the plate in the presence of sterile distilled water and sonicated at low output for 15 seconds. They were subsequently diluted to an A_{cop} = 0.1 and plated at dilutions of 10⁻³ and 10⁻⁴, in duplicate onto minimal plates containing glycerol as the carbon source, and so incubated at 30° C for 2-3 days. They were then replica-plated onto minimal plates to which 100 Lil of methanol was added in the vapor phase. After a 24-hour incubation at 30° C, it was apparent that 4% of the transformants were growing more slowly on methanol than the rest of the transformants. Five of the His Mut* isolates were examined by Southern analysis (Example V).

Example IV

Yeast DNA Miniprep

10° cells/ml were seeded in 5 ml YPD at 30° C overnight and then pelleted using a Damon IEC DPR600 clinical centrifuge at 3,000 rpm for 5 minutes. The pellet was resuspended in 0.5 ml of 1 M sorbiol, 0.1 ml 0.5 M EDTA, pH 8 and the sample transferred to a 1.5 ml microfuge tube. 0.02 ml of 2.5 mg/ml Zymolyase 100,000 (Miles Laboratories) was added, and the sample was incubated at 37° C for 60 minutes. The cells were pelleted using the microfuge for 1 minute at high speed, and resuspended in 0.5 ml of 50 mM Tris* Cl, 10° pH 7.4 and 20 mM EDTA. 0.05 ml of 10% SDS was added, the sample mixed, and incubated af 65° C for 30 minutes. 0.2 ml of 5 M potassium acetate, pH 5.2, was added and the sample was incubated on ice for 60 minutes. The sample was again spun in a microfuge at high speed for 5 minutes.

The supernatant was transferred to a fresh 1.5 ml microfuge tube and 1 volume of isopropanol at room temperature vas added. The sample was mixed and allowed to sit at room temperature for 5 minutes, then 5 spun very briefly (10 seconds) in a microfuge at high speed. The supernatant was poured off and the pellet air dried. After resuspending the pellet in 0.3 ml of 10 mM Tris* Cl, pH 7.4 and 1 mM EDTA, 15 µl of a 1 mg/ml solution of pancreatic RNase was added, and the sample was incubated at 37° C for 30 minutes. 0.03 ml of 3 M sodium acetate was added, the sample mixed, and 0.2 ml of isopropanol added. The sample was spun in a microfuge at high speed to pellet the DNA. The supernatant was then poured off, the pellet offield and resuspended in 0.1-0.3 ml of 10 mM Tris* Cl, pH 7.4 and 1 mM EDTA. (Note: Before using the DNA in a restriction digest, it may be necessary to spin the solution for 15 minutes at high speed in the microfuge to remove any insoluble material which may inhibit the digestion).

Example V

Strain Characterization

DNA was prepared from the transformed Pichia cells (Example III) and from untransformed host Pichia cells as described in Example IV, and digested with EcoRI. The samples were electrophoresed on 0.8% agarose gels, and Southern blots were performed (Manialis et al. 1982). The filters were hybridized with an AOXI specific probe or with a HIS4 specific probe to determine where integration had occurred. The site of integration was determined by comparing the spectrum of hybridization of a given transformant with the wild type strain. Any alteration in the size of the wild type band was evidence of integration at that locus. A summary of the Southern hybridizations and strain characterization for the strains chosen for further analysis is below.

Table I

Strain Name	Site of Integration	Vector Copy Number
G+HSA140S1	AOX1	one
G+HSA140S4	AOX1	two
G+HSA140S3	AOX1	>two
G-HSA140S1	AOX1	one

Fermentor Growth of HSA-Expressing Pichia Strains

Inocula were prepared from selective plates and grown overnight at 30 °C in buffered YNB containing 2% glycerol to an ODcas of 0.5-10.0. An aliquot of 5-50 ml of the overnight culture was added to a 2-little capacity termentor, and the repressed growth phase continued in 5X basal salts containing 5 ml/L of PTM, salts at 30 °C. The pH was maintained at 5.0 by the addition of 40% ((vi)) armonium hydroxide, and foaming was controlled by the addition of 5% ((vi)) Struktol artificam. Dissolved oxygen was maintained above 20% by increased aeration and agitation as needed. The temperature was maintained at about 30 °C. This batch growth phase continued for 20-30 hours until the glycerol was exhausted. The termentation was then continued in either a methanol-limited fed-batch mode for Mut* strains.

a. Mut fermentation; methanol-limited fed batch

Run 544: G+HSA140S1 (1 copy)

Run 557: G + HSA140S4 (2 copy)

Run 545: G + HSA140S3 (>2 copy)

In order to continue building cell density as well as to prevent the accumulation of excess ethanol in the fermentor, the AOX1 promoter was derepressed to allow the expression of a small amount of alcohol oxidase before induction by the addition of methanol. This derepression was achieved by growth under glycerol-limited conditions at about pH 5.0. Following exhaustion of the glycerol in the initial growth phase, a 10 50% (w/v) glycerol leaf (containing 12 ml/L of PTM, trace salts) was initiated at a rate of 8-16 ml/hour and continued until approximately 120-140 ml had been added. Full expression of the AOX1 promoter was then induced by the initiation of a methanol leed (100% McOH plus 12 ml/L PTM, trace salts) at 1 ml/hour. The methanol leed was maintained for several hours until the culture responded to methanol limitation. This response was expressed as a sudden rise in dissolved oxygen upon a brief cessation of the methanol feed. The methanol feed was then increased over an 8-12 hour period until a rate of 15. ml/hour was achieved. Fermentation was continued under these conditions for 82, 96 or 98 hours on methanol before the culture was harvested.

b. Mut- fermentation; methanol excess fed batch

The Mult fermentations were conducted as described for the Mult fermentations, except the MeOH feed was increased after 4 hours of 1 ml/hr feed to 3-4 ml/hr, to give a residual methanol concentration less than 0.5%.

Quantification of HSA Secreted into Growth Media

a. ELISA

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The ELISA procedure for human serum albumin requires the following reagents: Human Albumin (obtained 30 from Cappell, Organon Teknika), Goat anti-HSA antibody, peroxidase conjugated (obtained from Cappell, Organon Tecknika), and O-phenylenediamine (OPD), dichloride salt (obtained from Sigma, 10 mg/tablet). The HSA was reconstituted following the manufacturers directions. In this case (lot #26706) 3.0 ml of distilled water was added to the contents of the vial (the final concentration was 18.8 mg/ml), 29 aliquots of 100 µl each were labeled and quickfrozen.

35 16 aliquots were formed by diluting 100 μl of 18.8 mg/ml with 1.780 ml PBS (final 1.0 mg/ml). The 16 aliquots of 100 μl were labeled and quickfrozen.

100 more aliquots were formed by diluting 100 μl of 1 mg/ml with 9.9 ml PBS (final 10 μl). The 100 aliquots of 100 μl each, were then labled and quickfrozen. This dilution was used to begin the standard curve of dilutions. The goat anti-HSA reaspent was divided into 50 μl aliquots, labeled and quickfrozen. Goat anti-HSA conjugated to peroxidase was reconstituted by the addition of 2.0 mls of distilled water (final concentration 23 mg/ml). 50 μl of aliquots of the goat anti-HSA conjugate were then labeled and quickfrozen.

ELISA PROCEDURE:

Note: Use buffers at room temperatures only.

- Make up coating buffer immediately before use. Dilute goat anti-HSA antibody 1:500. Add 200 μl of this solution to each well. Parafilm tightly and incubate one hour at 37 °C.
- 2. Sharply flick contents of plate into sink. Wash 3 times with TBST. Wash 2 times with distilled water.
- 3. Add 200 µl blotto buffer to all wells. Parafilm tightly. Incubate overnight at 37 °C.
 - 4. Next morning, flick contents of wells into sink. Wash 3 times with TBST. Wash 2 times with distilled
 - 5. Add 100 µl of TBST to all wells.
 - 6. Dilute stock 10 µg/ml HSA standard with TBST.
 - S = stock from freezer = 10 µg/ml
 - SS = substack = 1:100 of S = 10,000 pg/100 µl
 - S-1 = 1:10 of SS 1,000 pg/100 µl
 - Dilute SS, 1:1 = 5,000 pg/100 µl

Dilute SS, 1:3.3 = 3,000 pg/100 μ I

Dilute SS, 1:5 = 2,000 pg/100 μ l Dilute S-1, 1:1 = 500 pg/100 μ l

Dilute S-1, 1:5 = 200 pg/100 µ1

Dilute 3.000 pg/100 µl 1:1 = 1,500 pg/100 µl

8. Add 100 µl sample dilutions and standard curve dilutions to each well.

Parafilm tightly and incubate 2 hours at 37 °C.

10. Wash five times with TBST. Wash two times with distilled water.

11. Dilute goat anti-HSA conjugate 1:2000 with blotto buffer. Incubate for two hours at room temperature in the dark.

12. Wash three times with TBST. Wash two times with distilled water.

13. Immediately before use: add one pellet of OPD to 3 mls of distilled water in a dark container. Pipet 21 mls of water into a 50 ml Falcon tube. Add 3 mls OPD solution to the Falcon tube, add 10 µl of 30% Hob.; and mix.

Add 200 µl of this solution to each well. Parafilm tightly and incubate 10 minutes in the dark. Stop the reaction by addition of 50 µl 4.5 M sulfuric acid.

14. Read on ELISA reader at 492 nm using filter 4.

b. Data

The level of HSA secreted from each of the strains, and other information pertinent to the fermentations, is provided in Table II:

Table II

г					101.0: 1		
- 1				Fermentation of I	HSA Strains		
	Run	Strain	Copy Number	Integration Site	Hours on MeOH	Cell Density (Wet) g/l	HSA in Broth g/l
30		G-HSA140S1 G+HSA140S1 G+HSA140S4 G+HSA140S3	1 1 2 >2	AOX1/Mut AOX1/Mut AOX1/Mut AOX1/Mut	98 96 82 96	445 415 450 353	0.971 0.964 0.754 0.185

Characterization of Recombinant Product

a. Gel analysis

Samples of fermentation broths of fermentation runs 537, 544 and 545 were withdrawn at different time points during the induction phase and analyzed by SDS gels and Coomassie blue staining, 521 but of a 10-fuld dilution of fermentor broth (equivalent to 05 bl) were applied to the gel. The relative intensity of the stained bands of rHSA (recombinant HSA) and the HSA standard confirmed the high (gram/liter) rHSA levels found by ELISA, In addition, the rHSA from all three fermentations and at all time points analyzed showed identical mobility with the HSA standard (69 Rd). The fact that the rHSA was the major protein species secreted by all HSA expression strains suggests high initial purity (PG9%) of the rHSA secreted into the growth medium. Another protein species which migrates at approximately 45 Kd could also be detected in increasing intensity with respect to fermentation time. This protein species may be produced in a secondary processing event during event during event during event during event during event during an proteopist growth product post-secretion by proteases secreted into the fermentation broth, or both, it is related to rHSA, as it was detectable on Western blots by HSA specific polyclonal antisera.

b. N-terminal sequence

Protein sequence of the N-terminal region of secreted rHSA was obtained on a dialysed sample of fermentor broth. The sequence was determined on an Applied Biosystems Model 470A protein sequencer. The phenythholydantion (PTH) derivatives of the amino acids were identified by high performance liquid

chromatography with an Applied Biosystems Model 120A analyzer. The results showed that the rHSA Nterminus is aspartic acid, consistent with the N-terminal amino acid of HSA. There does not appear to be any other precursor type of rHSA. The balance of the sequence determined was identical to the known sequence for HSA.

Example VI

Construction of HSA expression vector pHSA313

The pHSA313 vector was constructed to provide a vector with an exact linkage between the 3' end of the native AOX1 5'regulatory region (promoter) and the start codon of the HSA structural gene.

A. Creation of pHSA113bC1a

About 200 ng of pHSA113 (disclosed in European Patent Application 0 344 458 and shown in Figure 7) was digested at 37° 0 for 1 hour with 1 unit of Cali in 20 µl of REact 1 butfer. The digestion mixture was brought to 100 µl with water and extracted once with an equal volume of phenotichlorotrom:soamyl alcohol (25:24:1 V/V), followed by extracting the aqueous layer with an equal volume of chloroform:soamyl alcohol (24:1). The DNA in the aqueous phase was precipitated by adjusting the NACI concentration to 0.2 M and 20 adding 3 volumes of cold ethanol. The mixture was allowed to stand on ice (4°C) for 10 minutes and the DNA precipitate was collected by centrifugation for 30 minutes at 10,000 x g in a microfuge at 4°C. The DNA pellet was washed 2 times with 70% aqueous cold ethanol. The washed pellet was vacuum dried and dissoved in 10 µl water to which 2 µl of 10 x ligation butfer, 2 µl of 1 mg/ml BSA, 6 µl of water and 1 unit 1, bNA ligase were added. The mixture was incubated overnight at 4°C and a 10 µl aliquot was used to 25 transform E. coil DG75' (Maniatis, et al.) to obtain pHSA113aCla, which represents the deletion of HIS4 and 3'AQX1, along with small stretches of pBR322 sequences used to link these sequences. The deletion of the HIS4, 3'AQX1 and pBR322 sequences removes one of two Csp45I sites present in the pHSA113 vector. The remaining Csp45I site is in the AQX1 feedualtory region (promoter).

30 B. Creation of pXHSA113∆Cla

Digest 5 µg of pHSA113∆Cla for 1 hour at 37°C with 10 units of BstEll in 100 µl of REact 2 buffer. The digestion mixture was extracted with phenol and precipitated as detailed in step A. The DNA precipitate was dissolved in 100 µl of Csp45l buffer and digested at 37 °C for 2 hours in the presence of 10 units of Csp45l. 35 The digested DNA was then phenol extracted and precipitated as described in step A. The DNA precipitate was dissolved in 20 µl of water and 10 µl aliquots were loaded on 2 neighboring wells of a 0.9% agarose gel. Following electrophoresis, the gel portion corresponding to one of the lanes was stained and this was used to locate the position of the Csp45I-BstEll fragment of pHSA113∆Cla in the unstained lane. The gel portion containing the larger Csp45I-BstEII fragment of pHSA113ΔCla was excised from the gel. The gel portion containing the larger Csp45I-BstEII fragment was electroeluted into 500 µI of 5 mM EDTA, pH 8.0. The DNA solution was phenol extracted as detailed in step A and the DNA precipitate was dissolved in 100 μl water. The larger Csp45I-BstEll fragment was then ligated with the BstEll-Csp45I oligonucleotide linker described below. An aliquot (10 µI) of the Csp45I-BstEII fragments was ligated overnight at 4 °C with 20 ng of annealed linker olgonucleotides 5'-CGAAACG ATG AAG TGG (SEQ ID NO:8) and 5'-GTTACCCACT-45 TCATCGTTT (SEQ ID NO:9) in 20 μl ligase buffer containing 100 μg/ml BSA and 1 unit of T₄ DNA ligase. The ligation mixture was used to transform E. coli DG75' to obtain pXHSA113ΔCla. The pXHSA113ΔCla vector by virtue of the linker described above has an exact linkage between the 3' end of the native AOX1 5' regulatory region (promoter) and the HSA ATG start codon with no extraneous DNA sequences.

50 C. Creation of pHSA313

1 µg of pXHSA113ACIa was diposted for 4 hours at 37°C with Clal in 100 µl of REact 1 buffer. Following dipestion the reaction mixture was adjusted to alkaline phosphatase buffer conditions and treated with 10 units of call intestinal alkaline phosphatase in a 200 µl reaction volume for 30 minutes at 37°C. Phosphatase treatment was terminated by phenol extraction and the DNA was precipitated and dissolved in water at a concentration of approximately 10 ng/µl as described in step A and stored at 20°C.

1 µg of pA0807N (Figure 8, construction of which is described in European Patent Application 0 344 459) was digested for 4 hours at 37 °C with Pstl in 100 µl of REact 2 buffer. The digested DNA was

adjusted to alkaline phosphatase buffer conditions and treated with 10 units of calf intestinal alkaline phosphatase in a 200 µl reaction volume for 15 minutes at 55 °C. At the end of 15 minutes another 10 units of phosphatase was added and incubated for 15 minutes. Phosphatase treatment was terminated by phenol extraction and the DNA was precipitated as described in step A. DNA was digested for 4 hours at 37 °C with 5 units of Cali in 100 µl REAct 1 buffer containing 100 µl/ml BSA, followed by phenol extraction and precipitation of DNA as outlined in step A. The DNA precipitate was dissolved in water at a concentration of approximately 20 ng/µl. This Clal fragment contains the HIS4 gene and 3' AOX1 second insertable sequence.

Approximately 100 ng (10 µl) of Clal cleaved-phosphatased pXHSA113.Cla was mixed with approximately 80 ng of Pettl digested-phosphatased and Clal-cleaved pA0807N (4 µl), 4 ull of St ligase buffer, 2 ull of 1 mg/ml BSA and ligated overnight at 4 °C using 1 unit of 1, DNA ligase. The ligation mixture was used to transform E. coli DG75' to obtain pHSA313. The pHSA313 plasmid from this ligation comtains the complete pXHSA113AClas sequence liked to the HIS4 gene and the AOXI 3' second insertable sequence derived from pA0807N. The relative orientation of the components of the pHSA313 plasmid is the same as 15 that shown in Flaure 7 for obsamid pHSA113.

Example VII

Construction of 5' & 3' exact HSA expression plasmid pHSA413

The pHSA413 vector was constructed to provide a vector with an exact linkage between the 3' and of the AOXI 5' regulatory region and the start codon of the HSA structural gene as well as an exact linkage between the 5' end of the AOX1 3' termination sequence and the 3'end of the HSA structural gene.

25 A.. Creation of pXXHSA113∆Cla

1 μg of pXHSA113ΔCIa was digested for 4 hours at 37°C with 10 units of EcoRi in 100 μl REact 3 buffer. The digestion mixture was phenol extracted and DNA precipitated as detailed in Example VI. DNA precipitate was dissolved in 20 μl water and digested for 1 hour at 37°C with 20 units of Bsu38li in 100 μl 30 of Bsu38li buffer. The digestion mixture was phenol extracted, DNA precipitated and dissolved in 100 μl of water as detailed in Example VI. Approximately 100 ng of EcoRil and Bsu38li-bleaved DNA was mixed with 10 ng of annealed oligonucleotides 5'-TTAGGCTTATAAG (SEQ ID NO:10) and 5'-AATTCTTATAAGCC (SEQ ID NO:11) and ligated overnight at 4°C in 20 μl of T, DNA ligase buffer containing 100 μg/ml BSA and 10 units of T, DNA ligase. The ligation mixture was used to transform E. coli to obtain 30 pXHSA113ACIa. In this plasmid the sequence between Bsu38li and EcoRil (SEQ ID NO:12) present in DXHSA113ACIa. In this plasmid the sequence between Bsu38li and EcoRil (SEQ ID NO:12) present in DXHSA113ACIa.

Bsu36I

EcoRI

is replaced by 5'CC TTA GGC TTA TAA GAATTC (SEQ ID NO:13)

Bsu36I EcoRI

B. Creation of pHSA413

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1 μg of pXXHSA113ΔCla was digested for 4 hours at 37°C with Clal in 100 μl of REact 1 buffer. Following digestion the reaction mixture was adjusted to alkaline phosphatase buffer conditions and treated with 10 units of calf intestinal alkaline phosphatase in 200 μl reaction volume for 30 minutes at 37°C. Phosphatase treatment was terminated by phenol extraction and the DNA was precipitated and dissolved in

water at a concentration of approximately 10 ng/µl as described in step A and stored at -20 °C.

Approximately 100 ng (10 µl) of Clai cleaved-phosphatased pXHSA113Cla was mixed with approximately 80 ng (4 µl) of Patl digested phosphatased and Clai-cleaved pA8807N (see paragraph 2 in step 3 of Example VI), 4 µl of SX Tigase buffer, 2 µl of 1 mg/ml ESA and ligated overnight at 4 ° C using 1 unit of T. 5 DNA ligase. The ligation mixture was used to transform E. coil DG75' to obtain pHSA413. The pHSA413 plasmid from this ligation contains the complete pXMS413\tilde{\text{ACI}}\tilde{\text{Table Sequence}} and the AQXI 3' second insertable sequence derived from pA8607N. The relative orientation of the components of the pHSA413 plasmid is the same as that shown in Figure 7 for plasmid pHSA113.

10 Example VIII

Transformation of Pichia pastoris with pHSA313 and pHSA413

A. Vector preparation

About 10 μg each of pHSA313, pHSA413, and pA0907N (negative control) were digested for 12 hours at 37°C in 200 μid HS buffer with 50 units of Notl. The digested DNA samples were phenol extracted, precipitated as described in Example VI, dissolved lin 20 μl of CaS, and were then used for transformation of Pichila pastoris GS115. About 10 μg each of pHSA313, pHSA413, and pA0907N were also digested with 20 units of SS1 for 12 hours at 37°C in 200 μl of REat 2 buffer containing 100 μg/ml of BSA. The digested DNA samples were extracted with phenol, precipitated as described in Example VI and dissolved in 20 μl of CaS.

B. Cell Growth

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Pichia pastoris GS115 (NRRL Y-15851) was inoculated into about 10 ml of YPD medium and shake cultured at 30 °C for 12-20 hours. 100 ml of YPD medium was inoculated with a seed culture to give an ODsco of about 0.001. The medium was cultured in a shake flask at 30 °C for about 12-20 hours. The culture was harvested when the ODsco was about 0.2-0.3 by centrifugation at 1555 g for 5 minutes using a 30 Sorvall RBSC.

C. Preparation of Spheroplasts

The cells were washed in 10 ml of sterile water, and then centrifuged at 1500 g for 5 minutes.

(Centrifugation is performed after each cell wash at 1500 g for 5 minutes using a Sorroll RT6000B unless otherwise indicated.) The cells were washed once in 10 ml of freshly prepared SED, once in 10 ml of sterile 1M sorbitol, and finally resuspended in 10 ml of SCE buffer. 7.5 µt of 3 mg/ml Zymolyses (100,000 unitsg, obtained from Miles Laboratories) was added to the cell suspension. The cells were incubated at 30 °C for about 10 minutes. (A reduction of 60% in Ob₂₀₀ in 5% SDS can be utilized as a correct time marker.) The appropriates were washed in 10 ml of sterile 1 M sorbitol by centrifugation at 700 g for 5-10 minutes. 10 ml of sterile CaS was used as a final cell wash, and the cells were centrifuged again at 700 g for 5-10 minutes and then resuspended in 0.8 ml of CaS.

D. Transformation

Pichia pastoris GS115 cells were transformed with 10 Lig of linearized DNA (see step A) using the spheroplast transformation technique of Srewkinsha et al., Gene 59, 115-125 (1987). DNA samples were added (up to 20 µl volume) to 12 x 75 mm sterile polypropylene tubes. (DNA should be in a suitable buffer such as TE buffer or CaS), 100 µl of spheroplasts were added to each DNA sample and incubated at room temperature for about 20 minutes. I mil of PEG solution was added to each sample and incubated at room temperature for about 15 minutes and centrifuged at 700 g for 5-10 minutes. SOS (150 µl) was added to the pellet and incubated for 30 minutes at room temprature. Finally 850 µl of 1M sorbitol was added.

E. Regeneration of Spheroplasts

A bottom agarose layer of 20 ml of regeneration agar SDR was poured per plate at least 30 minutes before transformation samples were ready. In addition, 8 ml aliquots of regeneration agar were distributed to 15 ml conical bottom Corning tubes in a 45°C water bath during the period that transformation samples

were in SOS. Aliquots of 50 or 250 μ I of the transformed sample was added to the 8 ml aliquots of molten regeneration agar held at 45 °C and poured onto plates containing the solid 20 ml bottom agar layer. The plates were incubated at 30 °C for 3-5 days.

s F. Selection of Transformants

Transformants were selected for by culturing on SDR, a media lacking histidine. The colonies which given the absence of histidine were also screened for "methanol-slow" phenotype, indicating displacement of the AOX1 structural gene by the Notl DNA fragment) in the case of transformants obtained using Notl linearized vectors. Several transformed GS115 cells showing "methanol-normal" (those obtained with SSI linearized DNA) and methanol-slow were then cultured and assayed for the production of HSA.

Example IX

Methanol induced secretion of HSA in GS115/pHSA313, and GS115/pHSA413 Integrative Transformants

Pichia pastoris GS115 strains transformed with pHSA313 and pHSA413 were analysed for HSA secretion in shake tube cultures. Both methanol-slow and methanol-normal strains were used. In each case 36 independent clones were studied. Transformants obtained with pA0807N served as negative controls. A 20 protocol was developed to ensure efficient secretion and stable accumulation of HSA in the culture medium.

Cells were grown to saturation in 10 ml BMGR or BMGY, and were placed in 50 ml tubes (2-d days). The cells would be in the range of 10-20 A_{coo} units. The cells were harvested, the supernatant liquid was discarded, and then the pellet was resuspended in 2 ml of BMMR or BMMY. The tube was covered with a sterile gauze (cheese cloth) instead of a cap. The tube(s) were then returned to a 30 °C shaker. At the end of 2-2 days, the cells were pelleted, and the supernatant assayed for product. The pellets could be resuspended with fresh medium and returned to the shaker for renewed secretion. With Pichla+1SA strains, 10 ul of media supernatant was sufficient for analysis by SDS-PAGE followed by Coomassie statining. Under these conditions a single band of 67 kD corresponding to HSA was observed. There was no significant difference between the expression levels of 6S115(pHSA313 vs GS115(pHSA413 vs GS115(pHSA413 vs GS115(pHSA413) vs GS115(pHSA413 vs GS115(pHSA413) vs GS115(pHSA413 vs GS115(pHSA413) vs GS115(pHSA413 vs GS115(pHSA413 vs GS115(pHSA413) vs GS115(pHSA413 vs G

Example X

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Batch-Fed Fermentation of Mut- Pichia pastoris for Production of HSA

Pichia pastoris GS115:pHSA 413-6 was inoculated into a 20 liter Biolatifite fermentor with an 8.5 I working volume. The inoculum was prepared in the following manner: a culture was grown on a YM plate and then transferred to 100 ml YM broth in a shake flask and grown for about 24 hours. 50 mls of this culture was transferred to 1 liter of YM broth in a shake flask and also grown for about 24 hours. 51 liter of this was then transferred to 8.5 liters of fermentor medium in the Biolatifie fermentor. Fermentor medium consisted of Minimal salts + biotin + 5 percent glycerol. Batch growth conditions included the following: PH = 5.8 (controlled with NH₂), temperature = 30° C, and percent dissolved oxygen greater than 20 percent air saturation.

Glycerol exhaustion was complete after about 24 hours, at which time a slow melthanol feed was begun at a rate of 10-15 m/hr. The methanol concentration was monitored in the fermentor and the feed rate was adjusted to maintain a concentration of 0.5-0.9 percent of methanol in the broth.

Secreted HSA in the media was measured quantitaively by desitometry of Coomassie blue stained polyacrylamide gels containing SDS (SDS-PAGE). Areas were referenced to a series of known weights of authentic HSA run on the same SDS-PAGE gois. The data from these gels is included in Table III.

Table III

Production of HSA by Batch-Fed Fermentation													
Run	Strain	Run pH	Hrs. MeOH	Dry Cell Wt.	HSA in Broth g/l								
1	GS115:pHSA 413-6	5.79	101	ND	2.13								
2	GS115:pHSA 413-6	5.85	237	101	3.39								
3	GS115:pHSA 413-6	5.85	265	98.12	2.70								
4	GS115:pHSA 413-6	5.97	258	117	2.90								

SEQUENCE LISTING

(1) GENERAL INFORMATION:
(i) APPLICANT: Kotikanyadan Sreekrishna et al.
(ii) TITLE OF INVENTION: Expression of Human Serum Albumin in
Pichia pastoris
(iii) NUHBER OF SEQUENCES: 13
(iv) CORRESPONDENCE ADDRESS:
(A) ADDRESSEE: RICHMOND, PHILLIPS, HITCHCOCK & UMPHLETT
(B) STREET: P.O. Box 2443
(C) CITY: Bartlesville
(D) STATE: OK.
(E) COUNTRY: USA
(F) ZIP: 74005
(v) COMPUTER READABLE FORM:
(A) MEDIUM TYPE: Diskette
(B) COMPUTER: IBM PC
(C) OPERATING SYSTEM: PC-DOS/MS-DOS
(D) SOFTWARE: Display Write 4
(vi) CURRENT APPLICATION DATA:
(A) APPLICATION NUMBER:
(B) FILING DATE:
(C) CLASSIFICATION:
(viii) ATTORNEY/AGENT INFORMATION:
(A) NAME: Hal Brent Woodrow
(B) REGISTRATION NUMBER: 32,501
(C) REFERENCE/DOCKET-NUMBER: 32747

(ix) TELECOMMUNICATION INFORMATION:

(A) TELEPHONE: 1-918-661-0624

(2) INFORMATION FOR SEQ ID NO:1:

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- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 940 bp
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: Genomic DNA
- (ix) SEQUENCE DESCRIPTION: SEQ ID NO:1:

AGATCTAACA TCCAAAGACG AAAGGTTGAA TGAAACCTTT TTGCCATCCG ACATCCACAG GTCCATTCTC ACACATAAGT GCCAAACGCA ACAGGAGGGG ATACACTAGC AGCAGACCGT 120 TGCAAACGCA GGACCTCCAC TCCTCTTCTC CTCAACACCC ACTTTTGCCA TCGAAAAACC 180 AGCCCAGTTA TTGGGCTTGA TTGGAGCTCG CTCATTCCAA TTCCTTCTAT TAGGCTACTA 240 ACACCATGAC TTTATTAGCC TGTCTATCCT GGCCCCCCTG GCGAGGTTCA TGTTTGTTTA 300 TTTCCGAATG CAACAAGCTC CGCATTACAC CCGAACATCA CTCCAGATGA GGGCTTTCTG 360 AGTGTGGGGT CAAATAGTTT CATGTTCCCC AAATGGCCCA AAACTGACAG TTTAAACGCT 420 GTCTTGGAAC CTAATATGAC AAAAGCGTGA TCTCATCCAA GATGAACTAA GTTTGGTTCG 480 TTGAAATGCT AACGGCCAGT TGGTCAAAAA GAAACTTCCA AAAGTCGGCA TACCGTTTGT 540 CTTGTTTTGGT ATTGATTGAC GAATGCTCAA AAATAATCTC ATTAATGCTT AGCGCAGTCT 600 CTCTATCGCT TCTGAACCCC GGTGCACCTG TGCCGAAACG CAAATGGGGA AACACCCGCT 660 TTTTGGATGA TTATGCATTG TCTCCACATT GTATGCTTCC AAGATTCTGG TGGGAATACT 720 GCTGATAGCC TAACGTTCAT GATCAAAATT TAACTGTTCT AACCCCTACT TGACAGCAAT 780 ATATAAACAG AAGGAAGCTG CCCTGTCTTA AACCTTTTTT TTTATCATCA TTATTAGCTT 840 ACTTTCATAA TTGCGACTGG TTCCAATTGA CAAGCTTTTG ATTTTAACGA CTTTTAACGA 900 CAACTTGAGA AGATCAAAAA ACAACTAATT ATTCGAAACG

(3) INFORMATION FOR SEQ ID NO:2:

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- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 600 bp
 - (B) TYPE: nucleic acid (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: Genomic DNA
- (ix) SEQUENCE DESCRIPTION: SEQ ID NO:2:

TTCCGAGATT TAGTATACTT GCCCCTATAA GAAACGAAGG AAAGTAAACC CCATTCAATG TGAACAGAAA TCTTCCATTT ACCCCCCACT GGAGAGATCC 120 ATTTCAGCTT CCTTACCCCA GAAAAAAGAA ATTCGGACAA ATAGAACACT TTCTCAGCCA GCCCAAACGA ACAGATAATA ATTAAAGTCA TTCCATGCAC TCCCTTTAGC TGCCGTTCCA TCCCTTTGTT GAGCAACACC 240 AGGAAACTTA ACCGATACCT TGGAGAAATC TAAGGCGCGA 300 ATCGTTAGCC AGTACGAAAG CTTAGTGAAG GGTGTTCCGA TACCTTCTCC ACATTCAGTC 360 ATGAGTTTAG CCTAGATATC TCATGAAGAG ACGGAAACGG GCATTAAGGG TTAACCGCCA 420 ATAGATGGGC AGCTTTGTTA GTCCCCAGTT TAAAGTTTTT CTTTCCTATT CTTGTATCCT 480 AATTATAA AAGACAACAT ATAACAAGTT CGTTTTAACT TAAGACCAAA ACCAGTTACA 540 GAGTGACCGT TGTGTTTAAT ACACTAAAGT TCACTCTTAT CAAACTATCA AACATCAAAA 600 ACAAATTATA ACCCCTCTAA

(4) INFORMATION FOR SEQ ID NO:3:

(±)	SEQUENCE	CHARAC	TERIS	STICS:
	/ 1 × × × × × × × × × × × × × × × × × ×		1000	1 -

(A) LENGTH: 1830 bp (B) TYPE: nucleic acid (C) STRANDEDNESS: single

(D) TOPOLOGY: linear

(44) MOLECULE TYPE: Genomic DNA

	(11)	HOLE	CULE	TYPE:	Genor	ic D	NA								
	(ix)	SEQU	ENCE	DESCR	IPTION	: SE	Q ID	NO:	3:						
ATG Met	AAG Lys -35	TGG Trp	GTA Val	ACC Thr	TTT Phe	ATT Ile -30				Phe '					45
GCT Ala	TAT Tyr -20	TCC Ser	AGG Arg	GGT G1y	GTG Val	TTT Phe -15	CGT Arg	CGA Arg	GAT Asp	Ala	CAC His -10	AAG Lys	AGT Ser	GAG G1u	90
GTT Val	GCT Ala -5	CAT	CGG Arg	TTT Phe	AAA Lys	GAT Asp 1	TTG Leu	GGA Gly	GAA Glu	GAA Glu 5	AAT Asn	TTC Phe	AAA Lys	GCC Ala	135
TTG Leu 10	GTG Va 1	TTG Leu	ATT Ile		TTT Phe 15	GCT Ala	CAG G1n	TAT Tyr	CTT Leu	CAG Gln 20	CAG Gln	TGT Cys	CCA Pro	TTT Phe	180
GAA G1u 25	GAT Asp	CAT His	GTA Val		TTA Leu 30	GTG Val	AAT Asn	GAA Glu	GTA Val	ACT Thr 35	GAA Glu	TTT Phe	GCA A1a	AAA Lys	225
ACA Thr 40	TGT Cys	GTT Val	GCT Ala		GAG Glu 45	TCA Ser	GCT Ala	GAA Glu	AAT Asn	TGT Cys 50	GAC Asp	AAA Lys	TCA Ser	CTT Lue	270
CAT His 55	ACC Thr	CTT Leu	TTT Phe			AAA Lys	TTA Leu	TGC Cys	ACA Thr	GTT Val 65	GCA Ala	ACT	CTT	CGT	315
GAA G1u 70					ATG Met 75	GCT Ala	GAC Asp	TGC	TGT Cys	GCA Ala 80	AAA Lys	CAA G1n	GAA G1u	CCT	360
GAG Glu 85					TTC Phe 90	TTG Leu	GAA Gln	CAC	Lys	GAT Asp 95	GAC Asp	AAC Asn	CC/	AAC Asn	405
CTC Leu 100	Pro					CC#	GAG Glu	GTT Val	GA1	GTG Val	Met	TGC Cys	AC'	GCT r Ala	450
TTT Phe 115	His		AAT ASI			Thi	TTT Pbe	TTO Lev	Ly:	A AA/ 5 Lys 12:	в Туз	C TT/	TA'	T GAA r Glu	495

ATT GCC AGA AGA CAT CCT TAC TIT TAT GCC CGG GAA CTC CTT TIC 11e Ala Arg Arg His Pro Tyr Phe Tyr Ala Pro Glu Leu Leu Phe 130 TIT GCT AAA AGG TAT AAA GCT TAT AAA CTG CT GCT TIT ACA GAA TGT TGC CAA GCT Phe Ala Lys Arg Tyr Lys Ala Ala Phe Thr Glu Cys Cys Gln Ala 145 GCT GAT AAA AGG TAT AAA GCT GCC TGC TTT ACA GAA TGT GAC CAA CTT GGG Ala ASP Lys Ala Ala Ala 160 GAT GAA GGG AAG GT GCC TGC CTG CTG CAAA CAC AGA CTC GAT GAC TTG GGG AGT GAA GGG AAG GT GAA CTT CAAA TTG GAA TTG Asp Glu Gly Lys Vel 180 AGT CTC CAA AAA TT GAT GGA AGA GGT TTA AGA GAT TG GAA GTT Ser Leu Gln Lys Phe 190 GCT GC CTG AGC CAA AAA GAG GT TTA AAA GAG AGA GTT AAA GAA TTG Ala Arg Leu Ser Gln Arg Ala Arg Ale Phe Lys Ala Try Ala Val 195 TCC AAG TTA GTG ACA GAT TTC CAAA GCT GAG TTA GCA GAG ATA TAT ACC CAT GAA TTC CAAA AGT CT GAG GAA TTC CAT GGA GAT CTG CTT GAA AAT TTC CAAA AAT CTC AAT TTC CAAA AGT CAAG GAA TTC TGC AAG TAT ATC TGT GAA AAA ATT CAA GAT TGG GAA CTC GAG TTG GAC AGA TGT GAA GAA AAT CTC CTG GAA AAA AAT CTC CTG GAA AAA CTA GAA GAT CTG GAT AAA AAT CAA GAT TGG GAA CTT GCC AAG TAT ATC TGT GAA AAA CTA GAA TTC GAT GAA AAT CAAA AAT CAAA AAT CTC AAT GAG AGG GAA TTC GAG AAG TAT ATC TGT GAA AAA AAT CAA GAT TGG GAT CTC CTT GAA AAAA TTC CAAA AAT CTC AAG GAT CTC AAG GAA TTC GAG AAG TAT ATC TGT GAA AAA AAT CAA GAT TGG GAT CTC CATT GAC AAG TAT ATC TGT GAA AAA CTA GAT TGT GAA AAA AAT CAA GAT TGG GAT CTC CAGG TAT GCC AAG TAT ATC TGT GAA AAA AAT CAA GAT TGG GAT CTC CAGG AAA AAT CCC ACG GAA TTCC AGT AAA CTT GAG AAG TAT ATC TGT GAA AAA CTA GAT TGT GAA AAA CCT CTG TTG GAA AAA AAT CAA GAT TGG GAT CTC CAGT AAAA CTT GAG AAG TAT ATC TGT GAA AAA AAT CAA GAT TGG GAT CTC CAGG AAAA CTC AGG GAG ATG CTG GAA AAT GAT GAG AGG GAG CTT GCC AAG TGT GAA AAA AAT CAA GAT TGG GAT CTC CAGG AAAA CTC CAGG AAA AAT CCC AGG AAAA CCC TGC ATT GCC AAG TGG GAA AAT GAT GAG AGG GAG CTT GCC AAG TGG GAA AAT GAT GAG AGG GAT TGG AAAA AAT CCC AGG AAAA AAT CCC AGG AAAA AAA CCT CTG TGT GAA AAAA CCT CTG TGT GAA AAAA CCT CTG TGT GAT TGGC CAGG AAAAAA CCT CTG TGT GAT TGC CAGA CAGA											
Phe		I1e			Pro			Pro			540
ATIA ASP Lys Ala Ala Cys Leu Leu Pro Lys Leu Asp Glu Leu Arg 165 GAT GAA GGG AAG GTT TGG TCT GCC AAA CAC AGA CTC AAG TGT GCC 675 Asp Glu Gly Lys Val Ser Ser Ala Lys Gla Arg Leu Lys Cys Ala 175 AGT CTC CAA AAA TTT GGA GAA AGA GCT TTC AAA GCA TGG GCA GTA 720 190 GCT GCC CTG AGC CAG GAA AGA GCT TTC AAA GCA TGG GCA GTA 720 GCT Ala Arg Leu Ser Gla Cys		Phe			Lys			G1u			585
ASP GIU GIY Lys Val Ser Ser Ala Lys Gin Arg Leu Lys Cys Ala 180 AGT CIC CAA AAA TIT GGA GAA AGA GCT TIC AAA GCA TGG GCA GTA 720 190 GCT GC CTG AGC CAG AGA TIT CCC AAA GCT GAG TTI GCA GAA GTT CCC AAA GCT GAG GTA TGC TGC AGC GAG GTA 765 Ala Arg Leu Ser Gin Arg Phe Pro Lys Ala Giu Phe Ala Giu Val 205 TCC AAG TTA GTG ACA GAT CTT ACC AAA GTC CAC ACG GAA TGC TGC 220 CAT GGA GAT CTG CTT GAA AAT TT CCC AAA GTC CAC ACG GAA TGC TGC 220 CAT GGA GAT CTG CTT GAA AAT CTG CAC AGG GCG GAC CTT GCC GAG AGG TT GCA ACG GAT TGC CAC GAG AGG TT GCC CAC GAG AGG TGC TGC CAC GAG AGG TGC CAC GAG AGG TGC TGC CAC TGC CAC TGC CAC GAG AGG TGC TGC CAC GAG AGG TGC TGC CAC GAG AGG TGC TGC CAC TGC TGC CAC		Ala			Cys			Leu			630
Ser Leu Gin Lys Phe Gly Glu Arg Ala Phe Lys Ala Trp Ala Val 200 GCT GCC CTG AGC CAG AGA ATTT CCC AAA GCT GAG TTT GCA GAA GTT Ala Arg Leu Ser Gin Arg Phe Pro Lys Ala Glu Phe Ala Glu Val 205 TCC AAG TTA GTG AGA GAT CTT ACC AAA GTC CAC AGG GAA TGC TGC 220 CAT GGA GAT CTG CTT GAA AGT CTG CAC GAG GAT TGC TGC Ser Lys Leu Leu 235 CAT GGA GAT CTG CTT GAA AGT GCT GAG AGG GGG GAC CTT GGC His Gly Asp Leu Leu 235 AAG TAT ATC TGT GAA AAT CAA GAT TCG ATC TCC AGT AAA CTG AAG Lya Tyr 1le Cys Glu Asp Ser Ile Ser Ser Lys Leu Lys 255 GAA TGC TGT GAA AAA CCT CTG TGT GAA AAT CCA CAC GAT GAC AGG G1u Cys Cys Gru Lys Pro Leu Leu Glu Lys Ser His Cys Ile Ala 260 GAA GTG GAA AAT GAT GAG AGG GCG GAC CTT GCC G1u Val Glu Asn Asp GAT GAT GA GAT GCC ACC AGG GAT GAC AGG GCT GAT TTT GTT GAA AGT AGC AGG GCG GAC CTT GCC ALa GAT TGC TTG GAA AGT GAC AGG GCG GAC CTT GCC GCT GAT TTT GTT GAA AGT GAC AGG GCG GAC CTT GCC ALa GAT GAT GAT GAC AGG GCG GAC CTT GCC ALA GAT GCC GAA AAT GAT GAC AGG GCG GAC CTT GCC GCC AGT GAC AGG GAT GCC TCG TGC ATC GCC TCG AGT AGA CTG AGG GCT GAT TTT GTT GAA AGT GAC AGG GCG GAC TGT GCC ALA GAT GCT GAA AGT GAT GAC AGG GCG GAC CTT GCC ALA GAT GCT GAT TTT GTT GAA AGT GAC AGG GCG GAC GAT GCC TCG TGA GAC AGG GCG GAC GAT GCC TCG GAC AGG GAC GTT GCC GCC GAC GAC GAC GAC GAC GAC GAC GAC		Asp			Ser			Arg			675
Aia Arg Leu Ser Gln Arg Phe Pro Lys Ala Glu Phe Ala Glu Val 205 TCC AAG TTA GTG ACA GAT CTT ACC AAA GTC CAC ACG GAA TGC TGC Ser Lys Leu Val Thr 220 CAT GGA GAT CTG CTT GAA ACT CTG CTG GAT GAS ASS CTG GAA TGC TGC Cys Ala Asp Asp Arg Ala Asp Leu Ala 235 AAG TAT ATC TGT GAA AAT CAA GAT TCG ATC TCC AGT AAA CTG AAG Lys Tyr 1le Cys Glu Ass Glu Ass Glu Ass Gat GAC CTT GGC Glu Cys Cys Glu Lys Pro Leu Leu Leu Leu Leu Cys Ala Asp Asp Arg Ala Asp Leu Lys 256 GAA TGC TGT GAA AAA CCT CTG TGT GAA AAT CCA CTG CAC TG CAC ATG GGC Glu Cys Cys Glu Lys Pro Leu Leu Leu Leu Leu Glu Lys Ser His Cys Ile Ala 260 GAA GTG GAA AAT GAT GAA AGT CTG CTG TG TGC CAC TG CAC TG CAC Glu Val Glu Ass Asp Glu Hys Cys Cys Cys Glu Lys Ser His Cys Ile Ala 280 GCT GAT TTT GTT GAA AGT AGA CTG TGT CCA TG CAC TG CA		Ser			G1y			Lys			720
Ser Lys Leu Val Thr Asp Leu Thr Lys Val His Thr Glu Cys Cys	'	Ala			Arg			Glu			765
His Gly Asp Leu Leu Glu Cys Ala Asp Asp Arg Ala Asp Leu Ala 240 AAG TAT ATC TGT GAA AAT CAA GAT TGG ATC TGC AGT AAA CTG AAG 250 GAA TGC TGT GAA AAT CCT CTG TTG GAA AAA CCT CTG TTG GAA AAA CCT AAG CTG CTG ATT GCC AGT GAA CTG CTG CTG TTG GAA AAA CCT CTG TTG GAA AAA TGC ATG CTG ATT GCC ATT GCC ATT GCC ATT GCC ATT GCC CTG TTG CTG ATT GCC ATG CTG CTG TTG CTG ATT GCC ATG CTG TG TTG CTG ATT GCT ATT GCT CTG TU Val Glu Asn Asp Ctu Het Pro Ala Asp Leu Pro Ser Leu Ala 280 GCT GAT TTT GTT GAA AAG ATG TTT GCAAA AAC TAT GCT GAG ATG AAG ATG CTG CTG AAA AAC TAT GCT GAG ATG AAG ATG CTG CTG AAA AAC TAT GCT GAG AAG AAG AAG AAG AAG ATG CTG CTG AAA AAC TAT GCT GAG AAG AAG AAG AAG AAG AAG AAG AAG AA	141	Ser			Asp			His			810
Lys Tyr Ile Cys Glu Asn Gln Asp Ser Ile Ser Ser Lys Leu Lys 250 GAA TGC TGT GAA AAA CCT CTG TTG GAA AAA TGC CAC TGC ATT GGC 945 Glu Cys Cys Glu Lys Pro Leu Leu Glu Lys Ser His Cys Ile Ala 275 GAA GTG GAA AAT GAT GAC GAG ATG CCT GCT TGC TTG CTT ATT GGT 990 Glu Val Glu Asn Asp Glu Het Pro Ala Asp Leu Pro Ser Leu Ala 280 GCT GAT TTT GTT GAA AGT GTC TGC AAA AAC TAT GCT GAG 1035 Ala Asp Phe Val Glu Ser Lys Asp Val Cys Lys Asn Tyr Ala GTu 295 GCA AAG GAT GTC TGC TGC ATT TTG TAT GAA TAT GCT AGA 1080 Ala Lys Asp Val Phe Leu Gly Met Phe Leu Tyr Glu Tyr Ala GA)	His			G1u			Arg			855
G1u Cys Cys G1u Lys Pro Leu Leu G1u Lys Ser His Cys Ile Ala 265 GAA GTG GAA AAT GAT GAG ATG CTT GCT TGC TTG CTT TA GCT G1u Val G1u Asn Asp G1u Het Pro Ala Asp Leu Pro Ser Leu Ala 280 GCT GAT TTT GTT GAA AGT GAT GCT AAA AAC TAT GCT GAG Ala Asp Phe Val G1u Ser Lys Asp Val Cys Lys Asn Tyr Ala G1u 295 GCA AAG GAT GTC TTG GGC ATG TTT TTG TAT GAA TAT GCA AGA Ala Lys Asp Val Phe Leu Gly Met Phe Leu Tyr G1u Tyr Ala Arg		Lys			Asn			Ser	Ser		900
Glu Val Glu Asn Asp Glu Het Pro Ala Asp Leu Pro Ser Leu Ala 280 GCI GAT TIT GTI GAA AGT AAG GAT GTI TGC AAA AAC TAT GCI GAG Ala Asp Phe Val Glu Ser Lys Asp Val Cys Lys Asn Tyr Ala Glu 295 GCA AAG GAT GTC TGC GGC ATG TTI TGC TAT GAA TAT GCA AGA 1080 Ala Lys Asp Val Phe Leu Gly Met Phe Leu Tyr Glu Tyr Ala Arg		G1u			Pro			Ser	His		945
Ala Asp Phe Val Glu Ser Lys Asp Val Cys Lys Asn Tyr Ala Glu 300 305 GCA AAG GAT GTC TTC TG GGC ATG TTT TG TAT GAA TAT GCA AGA 1080 Ala Lys Asp Val Phe Leu Gly Met Phe Leu Tyr Glu Tyr Ala Arg	,	Glu			Glu			Lev	Pro		990
Ala Lys Asp Val Phe Leu Gly Met Phe Leu Tyr Glu Tyr Ala Arg	5	Ala			Ser			Lys	Ası		1035
		Ala			Leu			ту Туз	Gl:		1080

	AGG Arg 325	CAT His	CCT Pro	GAT Asp	TAC Tyr	TCT Ser 330			CTG Leu 335				1125
5	ACA Thr 340	TAT	GAA Glu	ACC Thr	ACT Thr	CTA Leu 345			GCC A1a 350				1170
10	CAT His 355	GAA Glu	TGC Cys	TAT Tyr	GCC Ala	AAA Lys 360			TTT Phe 365				1215
15	GAA Glu 370	GAG Glu	CCT Pro	CAG Gln	AAT Asn	TTA Leu 375			TGT Cys 380				1260
20	CAG G1n 385	CTT Leu	GGA Gly	GAG Glu	TAC Tyr	AAA Lys 390			CTA Leu 395				1305
	ACC Thr 400	AAG Lys	AAA Lys	GTA Val	CCC Pro	CAA Gln 405			ACT Thr 410				1350
25	TCA Ser 415	AGA Arg	AAC Asn	CTA Leu	GGA Gly	AAA Lys 420			TGT Cys 425				1395
30	GAA G1u 430	GCA A1a	AAA Lys	AGA Arg	ATG Met	CCC Pro 435				Leu		GTC Val	1440
35	CTG Leu 445	AAC Asn	CAG Gln	TTA Leu	TGT Cys	GTG Val 450				Pro		GAC Asp	1485
	AGA Arg 460	GTC Val	ACC Thr	AAA Lys	TGC Cys	TGC Cys 465				Asn		CCA Pro	1530
40	TGC Cys 475	TTT Phe	TCA Ser	GCT Ala	CTG Leu	GAA Glu 480				Val		GAG Glu	1575
45\	TTT Phe 490	AAT Asn	GCT Ala	GAA Glu	ACA Thr	TTC Phe 495				116		CTT Leu	1620
50	TCT Ser 505	GAG Glu	AAG Lys	GAG Glu	AGA Arg	CAA Gln 510				: Ala		GAG LG1u	1665

CTT Leu 520	GTG Val				CCC Pro 525						1710
GTT Val 535	ATG Het	GAT Asp			GCA Ala 540	GCT Ala					1755
GAC Asp 550	GAT Asp	AAG Lys	GAG G1u	ACC Thr	TGC Cys 555		GCC Ala				1800
GCT Ala 565	GCA Ala		CAA Gln	GCT Ala	GCC Ala 570			TAA			1830

(5) INFORMATION FOR SEO ID NO:4:

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- (1) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 36bp (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: Oligonucleotide
- (ix) SEQUENCE DESCRIPTION: SEQ ID NO:4:

CCCTCACACG CCTTTGAATT C ATG AAG TGG GTA ACC 36 Met Lys Trp Val Thr 1 . 5

- (6) INFORMATION FOR SEQ ID NO:5:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 39bp
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: Oligonucleotide
 - (ix) SEQUENCE DESCRIPTION: SEQ ID NO:5:

GCC TTA GGC TTA TAAGAATTCA GTTTAAAAGC ATCTCAG 39 Ala Leu Gly Leu 570

(7) INFORMATION FOR SEQ ID NO:6: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 18bp 5 ... (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: Screening Oligonucleotide (ix) SEQUENCE DESCRIPTION: SEQ ID NO:6: GCCTGGGAAT CC ATG AAG 18 Met Lys 1 (8) INFORMATION FOR SEQ ID NO:7: (1) SEQUENCE CHARACTERISTICS: (A) LENGTH: 18bp (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: Screening Oligonucleotide (ix) SEQUENCE DESCRIPTION: SEQ ID NO:7: TTA TAAGAATTCA GTTTA 18 Leu 573 (9) INFORMATION FOR SEQ ID NO:8: (1) SEQUENCE CHARACTERISTICS: (A) LENGTH: 16bp (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear (ii) MOLECULE TYPE: Oligonucleotide (ix) SEQUENCE DESCRIPTION: SEQ ID NO:8:

> CGAAACG ATG AAG TGG 16 Met Lys Trp

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	21 0 010 000 72
	(10) INFORMATION FOR SEQ ID NO:9: (1) SEQUENCE CHARACTERISTICS:
5	(A) LENGTH: 19bp (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear
	(ii) MOLECULE TYPE: Oligonucleotide
10	(ix) SEQUENCE DESCRIPTION: SEQ ID NO:9:
15	GTTACCCACT TCATCGTTT 19
	(11) INFORMATION FOR SEQ ID NO:10:
20	(1) SEQUENCE CHARACTERISTICS: (A) LENGTH: 13bp (B) TYPE: nucleic acid (C) STRANDELMESS: single (D) TOPOLOGY: linear
25	<pre>(ii) MOLECULE TYPE: Oligonucleotide (ix) SEQUENCE DESCRIPTION: SEQ ID NO:10:</pre>
	TTAGGCTTAT AAG 13
30	(12) INFORMATION FOR SEQ ID NO:11:

- (1) SEQUENCE CHARACTERISTICS:

 - (A) LENGTH: 14bp
 (B) TYPE: nucleic acid
 (C) STRANDEDNESS: single
 (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: Oligonucleotide
 - (ix) SEQUENCE DESCRIPTION: SEQ ID NO:11:

AATTCTTATA AGCC 14

(13)	INFORMATION	FOR	SEQ	ID	NO: 1	2:
------	-------------	-----	-----	----	-------	----

- (1) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 231bp
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: Linker Oligonucleotide
 - (ix) SEQUENCE DESCRIPTION: SEQ ID NO:12:
- CCTTAGGCTT ATAACATCTC TACATTTAAA AGCATCTCAG CCTACCATGA GAATAAGAGA 60
 - AAGAAAATGA AGATCAAAAG CTTATTCATC TGTGTTTTCT TTTTCGTTGG TGTAAAGCCA 120
 - ACACCCTGTC TAAAAAACAT AAATTTCTTT AATCATTTTG CCTCTTTTTC TCTGTGCTTC 180
- AATTAATAAA AAATGGAAAG AATCTAAAAA AAAAAAAAA AAAAGGAATT C 231
 - (14) INFORMATION FOR SEQ ID NO:13:
 - (1) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 20bp
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: Oligonucleotide
 - (ix) SEQUENCE DESCRIPTION: SEQ ID NO:13:
 - CCTTAGGCTT ATAAGAATTC 20

Claims

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- An expression cassette for the production of HSA in Pichia pastoris comprising

 a) a Pichia pastoris 5' regulatory region having a 5' end and a 3' end selected from a Pichia
 - a) a Pichia pastoris 5' regulatory region having a 5' end and a 3' end selected from a Pichia pastoris AOX1 5' regulatory region and a Pichia pastoris DAS1 5' regulatory region wherein the 3' end of the 5' regulatory region is operably linked to
 - b) a HSA structural gene encoding a HSA signal sequence and a mature HSA protein, having a 5' end and a 3' end wherein the HSA structural gene has an ATG start codon within no more than 11 decoxyribonucleotides of the 5' end of said HSA structural gene; operably linked to
 - c) a suitable 3' termination sequence.
- The expression cassette of claim 1 wherein the 5' regulatory region is selected from the AOX1 5' regulatory region and the DAS1 5' regulatory region from Pichia pastoris.
 - The expression cassette of claim 1 wherein the 3' termination sequence is isolated from a Pichia pastoris gene selected from an AOX1 gene, a DAS1 gene, a p40 gene and a HIS4 gene.
 - The expression cassette of claim 1 wherein the expression cassette is incorporated into a vector selected from circular plasmids and linear plasmids, the latter preferably being integrative site-specific vectors.

- The expression cassette of claim 1 wherein the expression cassette is incorporated into a vector comprising the following serial arrangement:
 - a) a first insertable DNA fragment;
 - b) at least one marker gene; and,

- c) a second insertable DNA fragment; wherein at least one expression cassette is incorporated either before or after the marker gene of component (b), and the first and second insertable DNA fragments employed are homologous with separate portions of the Pichia pastoris genome and the insertable fragments are in the same relative orientation as exist in the Pichia pastoris genome.
- 70 6. The expression cassette of claim 5 contained in said vector wherein the first insertable DNA fragment and the second insertable DNA fragment are obtained from the DNA sequences of a gene isolated from Pichia pastoris selected from an AOX1 gene, a pA9 gene, a DAS1 gene and an HIS4 gene.
 - The expression cassette of claim 5 contained in said vector wherein the marker gene is selected from a Pichia pastoris HIS4 gene, a Pichia pastoris ARG4 gene, a Saccharomyces cerevisiae SUC2 gene, a G418th gene of bacterial transposon Tn9031 and a G418th gene of bacterial transposon Tn9031.
 - 8. The expression cassette of claim 5 wherein said vector comprises
 - a) a first insertable DNA fragment which is an operable 5' regulatory region from the AOX1 gene being about one kilobase in length isolated from *Pichia pastoris* operably linked to
 - b) a HSA structural gene encoding a HSA signal sequence and a mature HSA protein having a 5' end and a 3' end wherein the HSA structural gene has a ATG start codon within no more than 8 decorviborucleotides of the 5' end of said HSA structural gene; operably linked to
 - c) the 3' termination sequence of the AOX1 gene isolated from Pichia pastoris; operably linked to
 - d) a marker gene which is the HIS4 gene isolated from Pichia pastoris; operably linked to
 - e) a second insertable DNA fragment which is about 0.65 kilobases of the AOX1 3' termination sequence.
 - The expression cassette of claim 8 wherein the HSA structural gene has a ATG start codon with either the deoxyribonucleotide AGGAATTC or no deoxyribonucleotide 5' of said ATG start codon.
 - 10. A Pichia pastoris cell transformed with an expression cassette according to any of claims 1 to 9.
- 11. The Pichia pastoris cell of claim 10 wherein the Pichia pastoris cell to be transformed is selected from Pichia pastoris GS195 (NRRL Y-1851), Pichia pastoris GS190 (NRRL Y-18014), Pichia pastoris (NRRL Y-11430) and Pichia pastoris (NRRL Y-11430), wherein Pichia pastoris (NRRL Y-11430), wherein Pichia pastoris GS 115 (NRRL Y-115851) is the most preferred Pichia pastoris GS 115 (NRRL Y-15851) is the most preferred Pichia pastoris GS
- A process for the secretion of HSA from transformed Pichia pastoris cells according to claim 10 or 11 comprising
 - a) transforming a *Pichia pastoris* cell with at least one vector having at least one expression cassette according to any of claims 1 to 9, and
 - b) culturing the resulting transformed Pichia pastoris cell under suitable conditions to obtain the production of HSA.

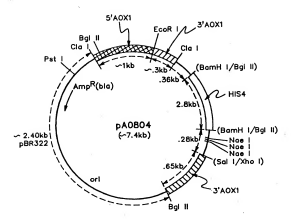


FIG. 1

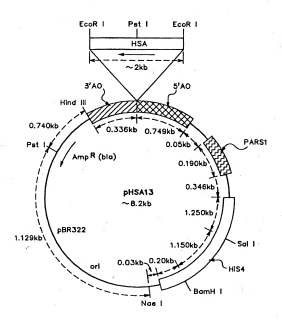


FIG. 2

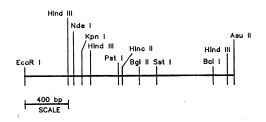


FIG. 3

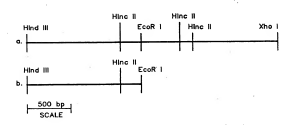


FIG. 4

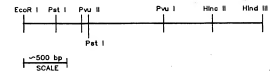


FIG. 5

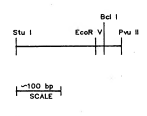
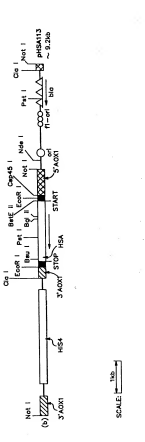


FIG. 6



.I.C. 7

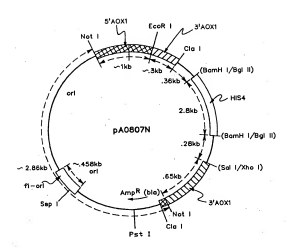


FIG. 8